

GenCore version 4.5
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OM nucleic - nucleic search, using sw model

Run on: September 4, 2002, 14:00:32 ; Search time 596.48 Seconds
(without alignments)
5684.861 Million cell updates/sec

Title: US-09-052-089a-8
Perfect score: 1975
Sequence: 1 GGACGACGAGTGGCGTGAGC.....CAAAAAAAAAAAAAAAAAAAAA 1975

Scoring table: IDENTITY_NUC
Gapop 10.0 , Gapext 1.0

Searched: 1736436 seqs, 858457221 residues

Total number of hits satisfying chosen parameters: 3472872

Minimum DB seq length: 0
Maximum DB seq length: 200000000

Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries

Database :
1: N_Geneseq_032802.*
2: /SID55/gcgdata/geneseq/geneseqn-emb1/NA1980.DAT.*
3: /SID55/gcgdata/geneseq/geneseqn-emb1/NA1981.DAT.*
4: /SID55/gcgdata/geneseq/geneseqn-emb1/NA1982.DAT.*
5: /SID55/gcgdata/geneseq/geneseqn-emb1/NA1983.DAT.*
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21: /SID55/gcgdata/geneseq/geneseqn-emb1/NA2000.DAT.*
22: /SID55/gcgdata/geneseq/geneseqn-emb1/NA2001A.DAT.*
23: /SID55/gcgdata/geneseq/geneseqn-emb1/NA2001B.DAT.*
24: /SID55/gcgdata/geneseq/geneseqn-emb1/NA2002.DAT.*

Pred. NO. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	1075.4	54.5	2065	20	AAx86754
2	1072.2	54.3	2065	19	AAV29062
3	100	5.1	148	22	AA527719
4	55.2	2.8	1506	23	AA575164
5	52.2	2.6	5154	23	AA584859
6	46.2	2.3	15377	13	AA025975
7	44.8	2.3	4246	22	AA560947
8	44.4	2.2	16442	18	AAx83006
9	44	2.2	744	21	AAc46364

10	44	2.2	746	21	AAc38464	Arabidopsis thalia
11	43.8	2.2	7453	22	AAI58369	Human polynucleoti
12	43.8	2.2	7501	22	AAI58370	Human polynucleoti
13	43.8	2.2	7741	22	AAI60155	Human polynucleoti
14	43.8	2.2	7741	22	AAI60156	Human polynucleoti
15	43.2	2.2	3229	22	AAK82155	Human immune/haema
16	42.8	2.2	390	23	AA569541	DNA encoding novel
17	42.8	2.2	3399	17	AAI05868	Chicken leucocytos
18	42.4	2.1	1080	24	ABI99537	Mouse ischemic co
19	41.6	2.1	2000	8	AAAT71065	Gene encoding Plas
20	41	2.1	6741	21	AAAI0595	Gene encoding a su
21	40.8	2.1	1994	21	AAE21149	Human low adenosin
22	40.8	2.1	1994	21	AAE21149	Human I-kappa-B kI
23	40.8	2.1	1994	21	AAE21149	Human adenosine re
24	40.8	2.1	2009	20	AA070513	Human RIP-associat
25	40.8	2.1	2034	20	AA070514	Human RIP-associat
26	40.8	2.1	8631	21	AAE21150	Human low adenosin
27	40.8	2.1	8631	21	AAE21150	Human adenosine re
28	40.4	2.0	3489	21	AAA30290	Kaposi's sarcoma-a
29	40.4	2.0	3489	22	AAE82901	Nucleotide sequenc
30	40.4	2.0	32207	20	AAV73805	KSHV LTR DNA (nucl
31	40.4	2.0	137507	19	AAV19941	KSHV long unique c
32	40.2	2.0	51259	18	AAx83007	Partial mouse WRN
33	40	2.0	941	21	AAE59944	Human secreted pro
34	40	2.0	5874	23	AA585420	DNA encoding novel
35	40	2.0	6413	22	AA546362	Tumour suppressor
36	40	2.0	10732	21	AAAI0584	Gene encoding a su
37	39.8	2.0	1931	23	ABU06323	Drosophila melanog
38	39.8	2.0	3931	23	ABU06322	Drosophila melanog
39	39.6	2.0	475	22	ABA58819	Human foetal liver
40	39.6	2.0	475	22	ABA27737	Probe #6203 for ge
41	39.6	2.0	475	22	AAK06973	Human brain expres
42	39.6	2.0	475	22	AAK32709	Human bone marrow
43	39.6	2.0	475	22	AAI38524	Probe #7210 used t
44	39.6	2.0	511	22	ABA71159	Human foetal liver
45	39.6	2.0	511	22	ABA37497	Probe #15963 for g

ALIGNMENTS

RESULT 1	
AAx86754	
ID	AAx86754 standard; cDNA: 2065 BP.
XX	
AC	AAx86754;
XX	
DT	27-OCT-1999 (first entry)
XX	
DE	cDNA 091-21A31 encoding a BRCA1 modulator protein.
XX	
KW	Modulator protein; BRCA1; tumour suppressor protein; breast cancer;
KW	ovarian cancer; cell growth; cell proliferation; ds.
XX	
OS	Homo sapiens.
XX	
FH	Key
FT	103..1512
FT	Location/Qualifiers
XX	/*tag= a
PN	US5948643-A.
XX	
PD	07-SEP-1999.
XX	
PF	13-AUG-1997; 97US-0968751.
XX	
PR	13-AUG-1997; 97US-0968751.
XX	
PA	(ONYX-) ONYX PHARM INC.
XX	
PI	Lingenfelter C, Polakis PG, Rubinfield B, Vuong TT;
XX	
DR	WPI; 1999-517952/43.

DR P-PSDB; AAY30149.

PT modulator proteins that bind to and modulate the activity of the
PT
BRCA1 tumour suppressor gene product, useful for the treatment of
PT ovarian and breast cancer

PS Claim 1; Fig 1; 35pp; English.

CC The present sequence encodes a modulator protein, that binds to and
CC modulate the activity of the BRCA1 gene product (BRCA1). The BRCA1
CC protein has been characterized as a tumour suppressor protein.
CC Alterations in the amino acid sequence of BRCA1 causes breast and ovarian
CC cancers by removing the controls on cell growth and proliferation.
CC Research has shown that different regions on the BRCA1 molecule have
CC different effects on cell growth and tumour suppression (e.g. full length
CC truncated BRCA1 has no effect on breast cancer cell growth but will
CC inhibit ovarian cancer cell growth). It has been suggested that different
CC host cell factors (e.g. proteins) interact with different regions of the
CC BRCA1 to control its function. The identification of these proteins
CC (e.g. BRCA1MP) will facilitate the development of novel diagnostic
CC methods and new therapeutics for identifying and treating cancers caused
CC by changes in the expression or activity of BRCA1.

Sequence 2065 BP; 561 A; 526 C; 561 G; 417 T; 0 other;

Query Match	54.5%	Score 1075.4;	DB 20;	Length 2065;
Best Local Similarly	75.1%	Pred. No. 1.7e-294;		
Matches 1460; Conservative	0;	Mismatches 426;	Indels 59;	Gaps 7;

Qy	81	TGATTCCTCCAGGAGCTCTCTGTAGATGACAGCCATCATGCGTATCTCCGCTCTGTGACATATCTG	1
Dp	72	Tgctgcctctggcccctctgaagtcacagccatcatgcctaccgcgtcctcgtgcacatctcgt	13
Oy	141	CTCCGACATCTTTCGATCATCTCCCGGAGCGGTGGCCATCCACGTGTGGCCACACTTTTCA	20
Dp	132	ctcgcacctctctgtactcactcccgcgagcgttgcgcgcacatccacatccacgtgcgcaccttcca	19
Oy	201	TTCTGCATTCCTTAATCCAGTCGTGTTGAGACAGACACCAAGTCGACCTTCGCCACAGTGTAG	26
Dp	192	cttgcagtcgctcaatctcagttgcttgaagacagcaccaagtcgaccttgcgcccacagttgcgc	25
Oy	261	AATCCAGCTGTGGCAAAAAGCATATTATTAACAACATTTCCTTACCCGCGCCAGGAGA	32
Dp	252	aatccaggtctggcaaaagaaacacatattacaataagctctctcttcttgccttgcgcccagggga	31
Oy	321	GGAGATGTTCTTGGATGACGAAATTTCTTAAGATGAACTGGACAGCGCTCAAGTCAAGCT	38
Dp	312	ggagaaatctcttgaatgaagaaatctcttaagaatgaacttgaacaaatgctcagaagccagct	37
Oy	381	TTCCGCAAAACACAGGGAGAAACGGGAGACGGCAGCGCCATTTCACACTCTACAGGACAC	44
Dp	372	ttccagagaagaacaagagaagaagacaagcagaagtcatactgacacactctgcgagatc	43
Oy	441	CCTGGAGACAGCGAATGCTACCGTGGAGTCCCTACAGACGCCCTTAAACAAGCAGAGAT	50
Dp	432	gctggaagaagcgaatgctacttggtaactctcttcgacagagccttggcaagggccagat	49
Oy	501	GCTGTGTTCCACCCGTAAGAAAACAGATGAATTCCTCGAGCAGCGGCGAGTATGAGACCA	56
Dp	492	gctgtgtcccaactctgaataaaagcagaatgaagtacttgaagcagcagcagatgagaccaa	55
Oy	561	ACAAAGCTTGGAGAGAGGCCACCGACTTCAGTGCAGAGTGAAGAAAACATGAGCAAAATTGA	62
Dp	552	acaaagcaaaagagagagccgcgctcgaagagaagaatgaagaagccatgagacagattga	61
Oy	621	GCTCTTACTCCAGAGCCAGCGCTTTCGAGGTGGAGAGATGATTTGAGACATGGGTGGG	68
Dp	612	gctcttactccaagggccagcgcccttgtaggttgaagagatgattccgaatacatgaggttgcggg	67
Oy	681	ACAAGTCAGCGGTGGAGCAGCTGCGTGTGTACTGCGGTGCTTCAAGAAAGATGTGAA	74
Dp	672	acacttcaagcgttgaacaacgcgcgtctgtactgtgtctctccaagaaaggttcaggaa	73

OY	741	TCGAGAGACAGCTGGGAAGCCACAGGGACACTGCTGACAGGTTGAAAGAGATTTCGT	800
Db	732	lctaaagaagcaagcaaggaagccctccagggagagtgagctctgacaagcttgaaggaatttctg	791
OY	801	GTCCTCTAGAGCAAGTTTGAAGACTCTCAACACTGTAGAGCTGGATTCAGGCCAATTTGAACT	860
Db	792	ttctccagaagaagcaagttgcagaacagtctcactcttgataatttgatacagccaagtctgaact	851
OY	861	GAGGTCAAGCCAGAGAGACTTTACAAAGTGTGACCAAGAGATACAGACTTAAGAAAGA	920
Db	852	gaatlcaagcccaagaagactctacagagtgcttgacaagaaatcatalgagcctctgaanaaaga	911
OY	921	GTCGATGATCTCTCAGAGGAACCTTGAGCCTGCTCC-GCGACCAATGAGAGGTCACAC	979
Db	912	g-ctaaagatgctgcgaagaaaccttgaaaccttcgacaccagtgccgaagtgcagctgcagcc	970
OY	980	GCTGTGTTTTGAGAGGCCCAAGCCCTGTGTGAGATGATGAAACCCGAGAGCTTCAACAGCCAC	1039
Db	971	gctcgtgttttagagtagcccaagcccccgtga--ggtgaatctgaaagctccgcgcgcat	1027
OY	1040	CTTTTCGGTGAATGAGATTGATCTCAATACCACTTTTATATGTAATACCTCTCAACCCAGA	1099
Db	1028	cccttcgttagatgatgatctcactcaatgctcactcttgatgtgatactcccccaagccgcgcg	1087
OY	1100	CCCTGTGCTCCACAGACTTGGCTCTCCCAAGAGACTGTGCTCGAGAGAGGACGCTTCCCA	1159
Db	1088	ctccagagctccccagcagtgttaactaagaaaaacttgccttagagaagtcaactccca	1147
OY	1160	TGCAGATGTCCTTCAGAGAGGTGCACAAAGCTTCCAGCCGGAATCCACGCTTCATCTGG	1219
Db	1148	ttcagagatgtcccccaagaagatactgaagaaagccccaagaaagatgccagactctacatcgg	1207
OY	1220	GTGGCAGGACATGTGTGAGAGACTGTGATGAGAACTGTGGCTGTGCTCTCCCTCTCTCA	1279
Db	1208	gtggccagagactgtgcagaggaagccagatgaaagaaacttgggtgccttccctaatllt	1267
OY	1280	TCGGAATGCTGTCTGGGTTCAGAAACAGGCCAACAGACACAGACAGAAATCCGAAGCA	1339
Db	1268	tcggaatgcatctcactcagccagaaacagcccaagggcccaagtcagagttccctctgca	1327
OY	1340	GCAAGATGTGTGAAGATATAGGCTTTGATGAGGCTTGAGAGAGCAAAATTCATCCAGC	1399
Db	1328	gcaaaagatgtgttaaggaacaggtccttcgaatgggtcgggtgcggaacaaatctacacgcg	1387
OY	1400	CTAAGGACACACACCATTAATTCGACAGATGGCTGTTAAGTTCAGAGGCCAAGAGTAACAGA	1459
Db	1388	ctactgacacaggtcatgatacgtcccatctgcgtttaagcccaaggaacaaagttaagcaga	1447
OY	1460	AAGTGAAGATTAAGACTGTGATGTTTCCTCCAGGCCCAAGCTGGATACCTTCTTATGTC	1519
Db	1448	gggtgaaggtgaagaacagtgctctctctctccagggccaagcttgacaaccttccctgtgt	1507
OY	1520	AG-----TGAAAGCTPACCAAGATGATGTTGTTTCAACATTAAGTGGGCCAACAC	1564
Db	1508	cgtgaagaacagtgatctgtgccaaatgycgaagacaacatgcccgtcaactgtgaatcaagga	1567
OY	1565	CTGGCTAACGGGAAGTGTTTTGTGGAAGATGGCTCTTGTGAGC-----	1608
Db	1568	ctgtccacagcgaggggttttctgtgacagagcccaacttcgggaacagccctgagtgtaag	1627
OY	1609	-----AGTCCAGACAGAGATGCCCGAANAACACACTTCCTGTGTTCACTG	1652
Db	1628	ggcagagaaacagtgtaggggtgtagtgtagaacccaagacatgctctctccctccacacc	1687
OY	1653	CGCCCTCTACAC--ACATGTGGGAAGCCACATGACCAATTTAGTGTCCATGACAGAGCC	1710
Db	1668	tgcccaactctacagactgtggagctgacatgacacagcccaactgatactcgtgaacaggtcc	1747
OY	1711	TACTTTCAGATTGCAAGGTTTTCCTTAATACCTACACACAGAGTGTGGCTTGAGACTCTTTGT	1770
Db	1748	tgctccagtttgcagagctcctctgtttatgaacatgataccaagtgttgcagagctcttccgg	1807

Qy	1771	TTTTNTAGAACAGGCTGCACATTACTCTTAAGTCATGGAGTGGAGATCTCTAGCA	1830
Db	1808	gctcgagaccacggtcactctgtgactctctctgtgaccag-----agtccttgagcc	1862
Qy	1831	GGCTGGAGGAGACCGGCTTGAACTCCTGCGCCGACGCTTATGCTTGAAATTGAG	1890
Db	1863	atctcagagcagcctcagcccaagcctctccacgaccttgaactgctcttcaggaagcct	1922
Qy	1891	GGTGAGGTGGTGTGATAGGGAAGGTTGGGGAAGTTTTCGTGTAATAATAAAAAGGATCT	1950
Db	1923	gggcgaagcagcggtggtgggaatgtgagatagatgcatgtgatgtatgtgagagatggaagatt	1982
Qy	1951	TTCTTCAAAAAAAAAAAAAAAAAA 1975	
Db	1983	tcatgtaaataataataataaaaaa 2007	
RESULT	2		
AAV29062			
ID	AAV29062	standard; cDNA; 2065 BP.	
XX	AAV29062;		
XX	28-AUG-1998	(first entry)	
XX	BRCA1	modulator protein 091-21A31 cDNA.	
DE	BRCA1	modulator protein; 091-21A31; breast cancer antigen 1;	
KW	tumour suppressor protein; diagnosis; therapy; human; ss.		
XX	Homo sapiens.		
XX	Key	Location/Qualifiers	
FT	CDS	103..1512	
FT		/+tag= a	
XX	WO9810066-A1.		
PN	12-MAR-1998.		
PD	06-AUG-1997;	97WO-US13944.	
PE	04-SEP-1996;	96US-0025601.	
XX	(ONLYX-)	ONLYX PHARM INC.	
PA	Liggenfelter C, Polakis P, Rubinfield B, Vuong TT;		
PI	WPI; 1998-193616/17.		
DR	P-PSDB; AAW37881.		
XX	Breast cancer antigen 1 modulator protein - useful for diagnosing		
PT	diseases involving unwanted cell growth, e.g. breast cancer, and for		
PT	producing therapeutics for treatment of such diseases		
XX	Claim 5; Fig 1; 73pp; English.		
XX	This cDNA clone, designated 091-21A31 (ATCC 98141), codes for		
CC	a 53 kDa BRCA1 modulator protein (see AAW37881) that binds to the		
CC	tumour suppressor gene product BRCA1, and which is characterised by		
CC	a zinc finger domain and a leucine zipper motif. 3 cDNA clones		
CC	(see also AAV29063 and AAV29064) coding for BRCA1 modulator proteins		
CC	(see AAW37881-83) were isolated from a HeLa cDNA library using a		
CC	yeast two-hybrid assay with a GAL4-BRCA1(8-1293) fusion as bait.		
CC	Vectors and host cells comprising the isolated nucleic acid		
CC	sequences are claimed, as well as a process for producing BRCA1		
CC	modulator protein by culturing these host cells. BRCA1 modulator		
CC	proteins and nucleic acids can be used to diagnose diseases		
CC	involving unwanted cell growth, e.g. breast cancer, and to identify		
CC	compounds that alter BRCA1 interaction with BRCA1 modulators for		
CC	the treatment of such diseases.		
XX	Sequence 2065 BP; 561 A; 528 C; 559 G; 417 T; 0 other;		

Query Match	54.3%	Score 1072.2	DB 19	Length 2065
Best Local Similarity	75.0%	Freq. No. 1.4e-293		
Matches 1458	Conservative	0	Mismatches 428	Indels 59
				Gaps
QY	81	TGGTTCCTCGGCGCTCTTGAGTGCAGACCATCATGCGCTATCTCTCTTGACATATCG	140	
DB	72	tgagcgcctcgggcccccttgaccacccaataatgctctcttgcctctcgtctcgtgacatctcg	131	
QY	141	CTCGCACTTCTTTCGATCATCTCCCGTAGCGTGGCTGCCATCATCTGTGGCCACACTTTTCA	200	
DB	132	ctccgactctcttgatactcccgagcgttgccgcgcacatccactcgcgcacacactcca	191	
QY	201	TCTGGAATGCCATATCCAGTGGTTTGAGACACACCAATGGAGACTGGCCACAGGTAG	260	
DB	132	cttgcagtgcccaatlcagtggtcttgagcagacccaagtgcgacctgcgccacagtgccg	251	
QY	261	AATCCAGGTTGGCAAAAAGACTATTTATAAACAACCTTTTCTTGACCTTGCCCGAGAGAA	320	
DB	252	aatccagtgctgcgcaaaagaccatatacaataaagtctctcttgccttgctgcgccagaga	311	
QY	321	GGAGAAATGTCCTTGATGCAGATTCTTAAAGATGACTGAGACGCGTCAAAGCTCAGCT	380	
DB	312	ggagaatctcttgatgctgcgaaatctctaaagaatgaactggaacaatgtaagagccagct	371	
QY	381	TTCCCGAAGACGAGGAGCAAAACGGGACGCGCAGGCGCATTTTCGACATCTCTGGGACAC	440	
DB	372	ttcccgagaagaagcaagggagaaacgagacagccagtgatcatatcgacaactctgcggatgc	431	
QY	441	CCTGGAGAGCAATGCTACCGTGGAGTCCCTACAGAAAGCGCTTTAAACAGGACAGAT	500	
DB	432	gctggaagaagcgaatgctactcgtgtatctctgcgacgagcctcttggaagccgagat	491	
QY	501	GCTGTCTTCCACCCCTGAAAAAACAGATGAATTTCTCGGAGACGGCGCAGAGATGAGACCA	560	
DB	492	gctgtgcttcacaactgaaaaagcagatgaaatgacttaagagcagcagcagatgaaagccaa	551	
QY	561	ACAACCTCGGAGGAGGCCACCACTCAAGTGCAGATGAAAGATGAAACCATGGACCAATTTA	620	
DB	552	acaagcaaaaggggggccccgcggtcgaagggcaagatgaaagacatgtagcagatgta	611	
QY	621	GCTCTACTCCAGAGCCAGCGTTCTGAGTGGAGAGAGATGATTCGAGACATGGGTGGG	680	
DB	612	gcttctactccagagccagcgccctgaagtgtagaggaatgataccgagacatggtgtg99	671	
QY	681	ACAGTCAGCGGTGGAGCACTGGGTGTATACGCGTGCCTTCACAAAGAGATATGAGA	740	
DB	672	acaagtcaggggtagaacacagctcgtgtagctgtgtagcttccaaagaagaagaaagaa	731	
QY	741	TCTGAAGAGAGCTCGGAAGGCCACAGGGGAGACTGCTGACAGGTGGAAGAGATTTGGT	800	
DB	732	tctaaagaaggcaggaagggctcaggggaaggttgctgtagcaagctgagaagatgtgt	791	
QY	801	GTCCTCTAGACACAAAGTTTAAAGACTCTCAACACTGAGCTGAGTCAAGCCAAAGTTAAGCT	860	
DB	792	tctctccaaagaagatgtgcagacagcttactcttgaaatttgatlcgaagccaagttagaact	851	
QY	861	GAGGTCAGCCCGAGAGAGACTTACAAAGTCTCTACACAGGAGATCACAGCGCTTAAGAAGAA	920	
DB	852	gaagtcagcgccagaaggaacttaacaggtgtctgaacaaggaatcatatgagctgtaaaagaa	911	
QY	921	GTCGTATGATCCTCCAGGGAACCTTGAGCCTGCGCTCC- GCGACCAATGAGCGGTTAGCC	979	
DB	912	g-ctaaagatgtctgcaggaagaaccttgaaaccttcacacaagatgycagatgagactgcac	970	
QY	980	GCGTGGTTTATAGAGCCCGCGCTGTGGAATGATGTAACCCGAGAGCTTACACACCCAC	1039	
DB	971	gectggttttagaagagccagccctctgta---gttgaatctgaaagctccgcgcgcacat	1027	
QY	1040	CCTTCGCGATGAGATTGATTCATATACCAACCTTTGATGTAAATACCCCTCCACCCAGA	1099	
DB	1028	cttcgcgtatataatctgcttcaatgtaactcttcatatcttgataatctccccagccggc	1087	

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QY 1100 CCTCTGGCTCCCGATTGGCTCCCAAGAGCTGCGCTGGAGAGGGCAGCTCCCA 1159
DB 1088 cctcagctcccgatgagtgatctactcgaaaaacttgcctgagagaagtcacactcccaa 1147
QY 1160 TGCAGATGCTCTCAAGAGAGTGCAAAAGTCTCCAAAGCCGAGTCCAGCTCTCACTGG 1219
DB 1148 ttcagatgctcccgaaagataltgcaaaagcccaaggaagagcccgctctcactcgg 1207
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DB 1328 gcaagatggtgtaaggaacagcttcgattggtcgtcggtcggaacaaattcaccagc 1387
QY 1400 CTAGGACACAAACCAATTCCGACAGTGCCTGTAACTCCAGGCCAAGAGTAACAGA 1459
DB 1388 ctactgacacagctcactgactcgccactgctgtaagcccaagcccaagtgtaagcaga 1447
QY 1460 AAGTGAGATTAAGACACTGTGAGTTCTGCCCTCCAGCCAGCTGAGTACCTTATATGC 1519
DB 1448 ggtgtaggggtgaagcagatgctctctctcccaagcccaagctgacacctcctcgtgt 1507
QY 1520 AG-----TGAAACGGTGACAGAGATGATTTTGCAATTAAGTGGGCCAAGAC 1564
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QY 1565 CTGGCTAACCGGAATGTTTTTGAAGATGGCTCCTCTTGAGAC----- 1608
DB 1568 ctgctcagcaggggtcttctgtagacagagcccccacttcgtagacagcctgagtgtaag 1627
QY 1609 -----ACTCCAGAGAGATGCCAGAAACACATCTCCTGTCTCACTG 1652
DB 1628 ggcagacaacagtgtaggggtgtagtgtagacacccagaaagcgtctctcctccacccc 1687
QY 1653 CGCCCTGCAC--ACACTGGGAGGCCACATGACAGTTACTGTTCCGATCAGAGGGCC 1710
DB 1688 tgcgccactctcactgagctggagctgacatgacccacactgactcctgtaagcagttcc 1747
QY 1711 TACTTCAGTTGACGGGTTTGTCTATAGCTACACAGCGGTGGCTGAGACTCCTTTGT 1770
DB 1748 tgcctcgttgcagagccctcgattatagcagatgcatcgatggtgctgagactccttcgg 1807
QY 1771 TTTTATAGACAGGCTCATTTGACTTGAAGTGAATGATGATGAGAGTGTGATCTATGCA 1830
DB 1808 gctcggagacacagtgctactgtagctgctcgtgtagcagc-----agtgctgagagc 1862
QY 1831 GGCTGGAGGACCTGCGCTTCACTGCTGCTCCAGCTTATGCTTGAATTAATGAG 1890
DB 1863 atctcagcagcctcagcccaagcttctactccttgccttgccttgcctcagtagcct 1922
QY 1891 GGTGAGTGTGATAGGAGAAAGTTGGGGAAGTTTCTGTGTAATAATAAAGGATCTT 1950
DB 1923 gggcccaagcagggctggggaagagatagcatggagatgtagagagatggaagattt 1982
QY 1951 TTCTTCAAAAAAAAAAAAAAAAA 1975
DB 1983 tcatgtataataataataaaaaa 2007
```

```
XX XX
DE DNA encoding novel signal transduction pathway protein, Seq ID 1379.
XX
KW Neuroprotective; cytoskeletal; dermatological; immunosuppressive; tumour;
KW antiinflammatory; anti-HIV; antibacterial; antiinflammatory; cancer;
KW immune system disorder; rheumatoid arthritis; inflammatory condition;
KW organ transplant rejection; infection; hepatitis C; blood disorder;
KW sickle cell anaemia; hyperproliferative disorder; Gaucher's disease;
KW neurodegenerative disorder; Alzheimer's disease; Parkinson's disease;
KW chromosomal abnormality; Down syndrome; ischaemia; renal disorder;
KW cardiovascular; respiratory; wound healing; endocrine; Addison's disease;
KW reproductive system; gastrointestinal; liver disorder; AIDS; ds;
KW acquired immune deficiency syndrome.
XX
OS Homo sapiens.
XX
PN WO200154733-A1.
PD
XX
XX 02-AUG-2001.
XX
PE 17-JAN-2001; 2001WO-US01312.
XX
PR 31-JAN-2000; 2000US-0179065.
PR 04-FEB-2000; 2000US-0180628.
PR 24-FEB-2000; 2000US-0184664.
PR 02-MAR-2000; 2000US-0186350.
PR 16-MAR-2000; 2000US-0189874.
PR 17-MAR-2000; 2000US-0190076.
PR 18-APR-2000; 2000US-0198123.
PR 19-MAY-2000; 2000US-0205515.
PR 07-JUN-2000; 2000US-0209467.
PR 28-JUN-2000; 2000US-0214886.
PR 30-JUN-2000; 2000US-0215135.
PR 07-JUL-2000; 2000US-0216647.
PR 07-JUL-2000; 2000US-0216880.
PR 11-JUL-2000; 2000US-0217487.
PR 11-JUL-2000; 2000US-0217496.
PR 14-JUL-2000; 2000US-0218290.
PR 26-JUL-2000; 2000US-0220963.
PR 26-JUL-2000; 2000US-0220964.
PR 14-AUG-2000; 2000US-0224518.
PR 14-AUG-2000; 2000US-0224519.
PR 14-AUG-2000; 2000US-0225213.
PR 14-AUG-2000; 2000US-0225214.
PR 14-AUG-2000; 2000US-0225265.
PR 14-AUG-2000; 2000US-0225267.
PR 14-AUG-2000; 2000US-0225268.
PR 14-AUG-2000; 2000US-0225270.
PR 14-AUG-2000; 2000US-0225447.
PR 14-AUG-2000; 2000US-0225757.
PR 14-AUG-2000; 2000US-0225758.
PR 14-AUG-2000; 2000US-0225782.
PR 14-AUG-2000; 2000US-0227009.
PR 30-AUG-2000; 2000US-0228924.
PR 01-SEP-2000; 2000US-0229287.
PR 01-SEP-2000; 2000US-0229343.
PR 01-SEP-2000; 2000US-0229344.
PR 01-SEP-2000; 2000US-0229345.
PR 05-SEP-2000; 2000US-0229509.
PR 05-SEP-2000; 2000US-0229513.
PR 05-SEP-2000; 2000US-0230437.
PR 06-SEP-2000; 2000US-0230438.
PR 08-SEP-2000; 2000US-0231242.
PR 08-SEP-2000; 2000US-0231243.
PR 08-SEP-2000; 2000US-0231244.
PR 08-SEP-2000; 2000US-0231413.
PR 08-SEP-2000; 2000US-0231414.
PR 08-SEP-2000; 2000US-0232080.
PR 08-SEP-2000; 2000US-0232081.
```


CC responsible for genetic disorders or other traits to assess biodiversity
CC and to produce other types of data and products dependent on DNA and
CC amino acid sequences. A564197-AAS94564 represent novel human
CC diagnostic coding sequences of the invention.
CC Note: The sequence data for this patent did not appear in the printed
CC specification, but was obtained in electronic format directly from WIPO
CC at [ftp.wipo.int/pub/published_pct_sequences](http://wipo.int/pub/published_pct_sequences).

XX
SQ Sequence 5154 BP; 1544 A; 1236 C; 1478 G; 896 T; 0 other;

Query Match 2.6%; Score 52.2; DB 23; Length 5154;
Best Local Similarity 48.5%; Pred. No. 0.00071;
Matches 175; Conservative 0; Mismatches 183; Indels 3; Gaps 1;

QY 485 TAAACAAGCCAGAGATGCTGTGTCACCCCTGAAAAAAGAGATGATGATCTCGAGACAGC 544
DB 494 tacagagctgagagagctgtccgaactcaaaacagatggtgaagaatgggctgc 553
QY 545 GGCAGAGTGAAGACCAAGCTCGGAGAGGCCACCGACTCAAGTGCANAGTGAATA 604
DB 554 tacagatggagccaacacctgagcagaggtgtgaaggtctgagggaaagctccaat 613
QY 605 CCATGAGACAAATTGAGCTCCTACTCCAGAGCCAGCTTCTGAGGTGAGAGATGATTC 664
DB 614 ccaggtcgaaacaacatcagccttgagctccttaagaagacaagaagcagactcc 673
QY 665 GAGCATGGGTGTGGGACAGTCACGGTGGAGCAGCTGGCTGT--GTACTGCTGTCCC 721
DB 674 aggagcagagagatctccgagagcagaggtgcagagagtcgagagcagagagac 733
QY 722 TCAGAAAGAGATGAGATGATCTGAAGAGAGCTCGAAGGCCACAGGGCAACTGGCTGACA 781
DB 734 tctgtgacaaacagagagcttcgagagcagcagaagacgtctcagagagcaggtgaga 793
QY 782 GGTGAAGAGGATTTGGTGTCTTAGAGAGCAAGTTGAGACACTCTCAACACTGAGCTGG 841
DB 794 ggcgtcgaaagcagagcagagctacgacaacagagagagctcgcgaaagagagag 853
QY 842 A 842
DB 854 a 854

RESULT 6
AAQ25975
ID AAQ25975 standard; DNA: 15377 BP.

XX AAQ25975;
XX
DT 08-JAN-1993 (first entry)
XX
DE MH mutant porcine ryanodine receptor cDNA.
XX
XX MH: RYR1; calcium release channel; sarcoplasmic reticulum;
KM transverse tubule; Pleitrain; Yorkshire; polymorphism; beta strand; ss.
XX
OS Synthetic.

XX
FH Key Location/Qualifiers
FT CDS 130..15237
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FT FT 207
FT FT variation /*tag= b
FT FT /*label= Polymorphic_site
FT FT 405
FT FT variation /*tag= c
FT FT /*label= Polymorphic_site
FT FT 438
FT FT variation /*tag= d
FT FT /*label= Polymorphic_site
FT FT 876
FT FT variation /*tag= e

FT FT /*label= Polymorphic_site
FT FT 1329
FT FT /*tag= f
FT FT /*label= Polymorphic_site
FT FT 1972
FT FT /*tag= g
FT FT /*label= MH_mutation
FT FT 2007
FT FT variation /*tag= h
FT FT /*label= Polymorphic_site
FT FT 4071
FT FT /*tag= i
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FT FT 4383
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FT FT 4462
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FT FT /*tag= n
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FT FT /*tag= p
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FT FT variation /*tag= q
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FT FT 14007
FT FT variation /*tag= w
FT FT /*label= Polymorphic_site
FT FT 15355..15360
FT FT /*tag= x

XX WO9211387-A.
XX PN
XX PD 09-JUL-1992.
XX
XX PE 20-DEC-1991; 91WO-CA00457.
XX PR 21-DEC-1990; 90GB-0027869.
XX PR 20-MAY-1991; 91GB-0010865.
XX PR 09-SEP-1991; 91GB-0019250.
XX
XX (UYGU-) UNIV GUELPH.
XX PA (UTOR) UNIV TORONTO INNOVATIONS FOUND.
XX PI MacLennan DH, O'Brien PJ;

DR	WP1: 1992-250106/30.
P	P-P5DB: AAR25450.
PT	Purified DNA mol. for diagnosis of porcine malignant hyperthermia
PT	- comprised DNA sequence encoding normal or mutant ryanodine
PT	receptor with specified endonuclease restriction map
XX	
PS	Disclosure; Fig 2; 96pp; English.
XX	
CC	The sequence given is the mutant pig ryanodine receptor (RYR1) gene
CC	from swine cDNA. The polymorphic sites were observed in comparisons
CC	of Plerstrain and Yorkshire breeds. There are 17 polymorphisms between
CC	the two breeds. The polymorphism at position 1972 causes a mutation
CC	from Arg to Cys and this is thought to be the molecular basis of
CC	porcine malignant hyperthermia (MH). This mutation lies within the
CC	region of RYR1 that is concerned with the binding of regulators of Ca ²⁺
CC	release channel gating. Analysis of surrounding sequences suggests
CC	that this mutation lies within a beta strand domain comprising roughly
CC	of amino acids 520 to 830. RYR1 is the calcium release channel of the
CC	sarcoplasmic reticulum and is a large protein which spans the gap
CC	between the transverse tubule and the sarcoplasmic reticulum. The
CC	channel is activated by ATP, calcium, caffeine, and micro-molar
CC	ryanodine. It is inhibited by ruthenium red, tetracaine, calmodulin,
CC	high Mg ²⁺ and ryanodine.
SQ	Sequence 15377 BP; 3197 A; 4630 C; 4755 G; 2774 T; 21 other;
XX	
Query Match	2.3%; Score 46.2; DB 13; Length 15377;
Best Local Similarity	51.2%; Pred. No. 0.065;
Matches 108; Conservative	0; Mismatches 103; Indels 0; Gaps 0;
OY	590 AGTCGACAGATGAACCACTGGAGCAAATTGACTCTTACTCCAGAGCAGCGTTCTGAGG 649
Db	
Db	5702 aggaagctgaaagaagatcttgtaagatgatctgagcccgaaagtattcaactgagaagaagg 5761
OY	650 TGCAGCAGATGATTTCCGACACATGSGGTGTGGCACAGTCAAGCGGTGGACAGCTGGCTGTGT 709
Db	
OY	710 ACTGCGTGTCCCTCAAGAAAGAGTATGAGAAATCTGAGAAGAGACCTGGAAGGCCACAGGG 769
Db	
OY	770 AACCTGCTGACAGGTTTGAAAGAAAGATTGGT 800
Db	
OY	5882 aaaagaagaatacttggagaaggaggtgcgt 5912
RESULT	7
AAS60947	ID AAS60947 standard; CDNA: 4246 BP.
XX	AAS60947;
XX	29-JAN-2002 (first entry)
DE	Human cancer agent-resistance marker #606.
XX	
KW	Human; cancer cell marker; TAXOL; cytosolic; tumour; carcinoma;
KW	squamous cell carcinoma; sarcoma; fibrosarcoma; leukemia;
KW	lymphocytic leukaemia; lymphoma; plasmacytoma; reticulum cell sarcoma;
XX	Hodgkin's disease; glioma; ss.
OS	Homo sapiens.
PN	WO200179556-A2.
PD	25-OCT-2001.
PF	13-APR-2001; 2001WO-US12132.
PR	14-APR-2000; 2000US-197538P.

(MILL-) MILLENNIUM PREDICTIVE MEDICINE INC.
Lillie J., Brown JL, Bolt A, Van Huffel C;
WPI; 2001-602933/68.

Novel nucleic acid, used as a marker to determine the effectiveness of
using TAXOL to treat cancer cell growth in individuals -

Claim 1; Page 450-451; 527pp; English.

The invention relates to 1046 novel nucleic acids which are used as
markers for determining the sensitivity of a cancer cell to the
anticancer agent TAXOL. Cancer cells can be treated with TAXOL when
they are shown to express one of the 242 sensitivity markers or the
cells are shown not to express one of the 804 resistance markers.
The methods can be used to determine the effectiveness of TAXOL
in the treatment of cancer cell growth in an individual. The markers
can be used as targets in developing anti-cancer agents such as
chemotherapeutic compounds. The markers can also be used as targets in
developing treatments for cancer, particularly those cancers which
display resistance to agents and exhibit expression of the markers. The
anticancer agents developed by the novel method can be used to treat
cancer. Probes based on the markers can be used to detect transcripts or
genomic sequences corresponding to the markers, in the identification of
cells or tissues which mis-express the protein. Cancers which may
be targeted include carcinoma (e.g. squamous cell carcinoma),
sarcoma (e.g. fibrosarcoma) leukaemia (e.g. lymphocytic leukaemia),
lymphoma, plasmacytoma, reticulum cell sarcoma, Hodgkin's disease and
tumours (e.g. glioma). The present sequence is one of the 1046
novel cancer cell markers.

Sequence 4246 BP; 1217 A; 1019 C; 1183 G; 827 T; 0 other;

[illegible]

DE Partial mouse WRN genomic sequence #2.
XX
XX Mouse; WRN; Werner's syndrome; detection; diagnosis; autosomal;
KM recessive disorder; phenotype: ss.
XX
OS Mus musculus.
XX
XX WO9724435-A1.
XX
XX 10-JUL-1997.
XX
XX 30-DEC-1996; 96WO-US20785.
XX
XX 12-APR-1996; 96US-0632175.
XX 29-DEC-1995; 95US-0009409.
XX 29-DEC-1995; 95US-0580539.
XX 30-JAN-1996; 96US-0010835.
XX 30-JAN-1996; 96US-0594242.
XX
XX (DARW-) DARWIN MOLECULAR CORP.
XX (OSHI/) OSHIMA J.
XX
XX Fu Y, Mulligan J, Oshima J, Schellenberg GD, Yu C;
XX
XX WPI: 1997-363671/33.
XX
XX Isolated nucleic acid molecule encoding the WRN gene product -
PT useful for detection and treatment of Werner's syndrome, and related
PT diseases
XX
XX
XX Claim 1; Fig 7; 153pp; English.
XX
XX This sequence represents a fragment of the genomic sequence containing
CC the coding region for the mouse WRN gene (AAK3004). The corresponding
CC human gene (AAK3001) encodes a protein related to Werner's syndrome.
CC The products can be used for the detection and treatment of Werner's
CC syndrome (WS), an autosomal recessive disorder with a complex phenotype,
CC as well as related diseases.
XX
XX
SQ Sequence 16442 BP; 4392 A; 2975 C; 3408 G; 5665 T; 2 other:

Query Match 2.2%; Score 44.4; DB 18; Length 16442;
Best Local Similarity 46.2%; Pred. No. 0.22;
Matches 147; Conservative 0; Mismatches 171; Indels 0; Gaps 0;

QY 535 CTGAGACGCGCGAGATGAGACCAACAGAGCTCGGAGAGGCCACCGACTCAAGTGC 594
DB 16438 CAGGAGGAGGAGCAGAGCAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAGAG 16379
QY 595 AAGATGAANAACCATGAGCAATGAGCTCTACTCCAGAGCCAGCGTTGAGAGTGAG 654
DB 16378 GAGAGGAGGAGCAG 16319
QY 655 GAGATGATTCGAGCATGCTGTGGAGACAGCGGAGAGAGCTGCTGTGTAAGTGC 714
DB 16318 CAGAGAGGAGCAG 16259
QY 715 GTGTCCCTCAAGAAAGATATGAAATCTGAAGAAAGCTCGAAGGCCACAGGGAAGCTG 774
DB 16258 CAGAGAGGAGCAG 16199
QY 775 GCTACACAGTTGAAAGAGATTTGTGTCTCTAGAGCAAGTTGAAGACTCTCAACTCT 834
DB 16198 AAGGAG 16139
QY 835 GAGCTGATCAGGCCAAG 852
DB 16138 AAGAGAGAGAGAGAGAGAG 16121

RESULT 9
AAC46364

ID AAC46364 standard; DNA: 744 BP.
XX
XX AAC46364;
AC
XX 18-OCT-2000 (first entry)
DT
XX
XX Arabidopsis thaliana DNA fragment SEQ ID NO: 49870.
DE
XX
XX Hybridisation assay; genetic mapping; gene expression control;
KM protein identification; signal transduction pathway;
KM metabolic pathway; promoter; termination sequence; ss.
XX
XX Arabidopsis thaliana.
XX
XX EPI033405-A2.
XX
XX 06-SEP-2000.
XX
XX
XX 25-FEB-2000; 2000EP-0301439.
XX
XX 25-FEB-1999; 99US-0121825.
XX 03-MAR-1999; 99US-0123180.
XX 09-MAR-1999; 99US-0123548.
XX 23-MAR-1999; 99US-0125788.
XX 25-MAR-1999; 99US-0126264.
XX 29-MAR-1999; 99US-0126785.
XX 01-APR-1999; 99US-0127462.
XX 06-APR-1999; 99US-0128234.
XX 08-APR-1999; 99US-0128714.
XX 16-APR-1999; 99US-0129845.
XX 19-APR-1999; 99US-0130077.
XX 21-APR-1999; 99US-0130449.
XX 23-APR-1999; 99US-0130510.
XX 23-APR-1999; 99US-0130891.
XX 28-APR-1999; 99US-0131449.
XX 30-APR-1999; 99US-0132048.
XX 30-APR-1999; 99US-0132407.
XX 04-MAY-1999; 99US-0132484.
XX 05-MAY-1999; 99US-0132485.
XX 06-MAY-1999; 99US-0132486.
XX 06-MAY-1999; 99US-0132487.
XX 07-MAY-1999; 99US-0132863.
XX 11-MAY-1999; 99US-0134256.
XX 14-MAY-1999; 99US-0134218.
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XX 18-MAY-1999; 99US-0134768.
XX 19-MAY-1999; 99US-0134941.
XX 20-MAY-1999; 99US-0135124.
XX 21-MAY-1999; 99US-0135353.
XX 24-MAY-1999; 99US-0135629.
XX 25-MAY-1999; 99US-0136021.
XX 27-MAY-1999; 99US-0136392.
XX 28-MAY-1999; 99US-0136782.
XX 01-JUN-1999; 99US-0137222.
XX 03-JUN-1999; 99US-0137528.
XX 04-JUN-1999; 99US-0137502.
XX 07-JUN-1999; 99US-0137724.
XX 08-JUN-1999; 99US-0138094.
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XX 14-JUN-1999; 99US-0139119.
XX 16-JUN-1999; 99US-0139452.
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XX 17-JUN-1999; 99US-0139492.
XX 18-JUN-1999; 99US-0139454.
XX 18-JUN-1999; 99US-0139455.
XX 18-JUN-1999; 99US-0139456.
XX 18-JUN-1999; 99US-0139457.
XX 18-JUN-1999; 99US-0139458.
XX 18-JUN-1999; 99US-0139459.
XX 18-JUN-1999; 99US-0139460.

Db 348 tcaagaatgagcttgatgatctca 371

RESULT 10

ID AAC38464 standard; DNA; 746 BP.

XX

AC AAC38464;

XX

DT 17-OCT-2000 (first entry)

XX

DE Arabidopsis thaliana DNA fragment SEQ ID NO: 21066.

XX

KM Hybridisation assay; genetic mapping; gene expression control;

KW protein identification; signal transduction pathway;

KW metabolic pathway; promoter; termination sequence; ss.

XX

OS Arabidopsis thaliana.

XX

PN EPI033405-A2.

XX

PD 06-SEP-2000.

XX

PF 25-FEB-2000; 2000EP-0301439.

XX

PR 25-FEB-1999; 99US-0121825.

PR 05-MAR-1999; 99US-0123180.

PR 09-MAR-1999; 99US-0123548.

PR 23-MAR-1999; 99US-0125788.

PR 25-MAR-1999; 99US-0126264.

PR 29-MAR-1999; 99US-0126785.

PR 01-APR-1999; 99US-0127462.

PR 06-APR-1999; 99US-0128234.

PR 08-APR-1999; 99US-0128714.

PR 16-APR-1999; 99US-0129845.

PR 19-APR-1999; 99US-0130077.

PR 21-APR-1999; 99US-0130449.

PR 23-APR-1999; 99US-0130510.

PR 23-APR-1999; 99US-0130891.

PR 28-APR-1999; 99US-0131449.

PR 30-APR-1999; 99US-0132048.

PR 30-APR-1999; 99US-0132407.

PR 04-MAY-1999; 99US-0132484.

PR 05-MAY-1999; 99US-0132485.

PR 06-MAY-1999; 99US-0132486.

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PR 07-MAY-1999; 99US-0132863.

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PR 18-MAY-1999; 99US-0134768.

PR 19-MAY-1999; 99US-0134941.

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PR 21-MAY-1999; 99US-0135353.

PR 24-MAY-1999; 99US-0135629.

PR 25-MAY-1999; 99US-0136021.

PR 27-MAY-1999; 99US-0136392.

PR 28-MAY-1999; 99US-0136782.

PR 01-JUN-1999; 99US-0137222.

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PR 18-JUN-1999; 99US-0139462.

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PR 22-JUN-1999; 99US-0139899.

PR 23-JUN-1999; 99US-0140353.

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PR 24-JUN-1999; 99US-0140695.

PR 28-JUN-1999; 99US-0140823.

PR 29-JUN-1999; 99US-0140991.

PR 30-JUN-1999; 99US-0141287.

PR 01-JUL-1999; 99US-0141842.

PR 01-JUL-1999; 99US-0142154.

PR 02-JUL-1999; 99US-0142055.

PR 06-JUL-1999; 99US-0142350.

PR 08-JUL-1999; 99US-0142803.

PR 09-JUL-1999; 99US-0142920.

PR 12-JUL-1999; 99US-0142977.

PR 13-JUL-1999; 99US-0143542.

PR 14-JUL-1999; 99US-0143624.

PR 15-JUL-1999; 99US-0144005.

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PR 16-JUL-1999; 99US-0144086.

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PR 19-JUL-1999; 99US-0144333.

PR 19-JUL-1999; 99US-0144334.

PR 20-JUL-1999; 99US-0144352.

PR 20-JUL-1999; 99US-0144632.

PR 20-JUL-1999; 99US-0144884.

PR 21-JUL-1999; 99US-0144814.

PR 21-JUL-1999; 99US-0145086.

PR 21-JUL-1999; 99US-0145088.

PR 22-JUL-1999; 99US-0145085.

PR 22-JUL-1999; 99US-0145087.

PR 22-JUL-1999; 99US-0145089.

PR 22-JUL-1999; 99US-0145192.

PR 23-JUL-1999; 99US-0145145.

PR 23-JUL-1999; 99US-0145218.

PR 23-JUL-1999; 99US-0145224.

PR 26-JUL-1999; 99US-0145276.

PR 27-JUL-1999; 99US-0145913.

PR 27-JUL-1999; 99US-0145918.

PR 27-JUL-1999; 99US-0145919.

PR 28-JUL-1999; 99US-0145951.

PR 02-AUG-1999; 99US-0146386.

PR 02-AUG-1999; 99US-0146388.

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PR 03-AUG-1999; 99US-0147038.

PR 04-AUG-1999; 99US-0147204.

PR 04-AUG-1999; 99US-0147302.

PR 05-AUG-1999; 99US-0147192.

PR 05-AUG-1999; 99US-0147260.

PR 06-AUG-1999; 99US-0147303.

PR 06-AUG-1999; 99US-0147416.

PR 09-AUG-1999; 99US-0147493.

PR 09-AUG-1999; 99US-0147935.

PR 10-AUG-1999; 99US-0148171.

PR 11-AUG-1999; 99US-0148319.

PR 12-AUG-1999; 99US-0148341.

PR 13-AUG-1999; 99US-0148565.

PR 13-AUG-1999; 99US-0148664.

PR 16-AUG-1999; 99US-0149368.

PR 17-AUG-1999; 99US-0149175.

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PR 18-AUG-1999; 99US-0149426.
PR 20-AUG-1999; 99US-0149722.
PR 20-AUG-1999; 99US-0149723.
PR 20-AUG-1999; 99US-0149929.
PR 23-AUG-1999; 99US-0149902.
PR 23-AUG-1999; 99US-0149920.
PR 25-AUG-1999; 99US-0150566.
PR 26-AUG-1999; 99US-0150884.
PR 27-AUG-1999; 99US-0151065.
PR 27-AUG-1999; 99US-0151066.
PR 27-AUG-1999; 99US-0151080.
PR 30-AUG-1999; 99US-0151303.
PR 31-AUG-1999; 99US-0151438.
PR 01-SEP-1999; 99US-0151930.
PR 07-SEP-1999; 99US-0152363.
PR 10-SEP-1999; 99US-0153070.
PR 13-SEP-1999; 99US-0153758.
PR 15-SEP-1999; 99US-0154018.
PR 16-SEP-1999; 99US-0154039.
PR 20-SEP-1999; 99US-0154779.
PR 22-SEP-1999; 99US-0155139.
PR 23-SEP-1999; 99US-0155486.
PR 24-SEP-1999; 99US-0155659.
PR 28-SEP-1999; 99US-0156458.
PR 29-SEP-1999; 99US-0156596.
PR 04-OCT-1999; 99US-0157117.
PR 05-OCT-1999; 99US-0157753.
PR 06-OCT-1999; 99US-0157865.
PR 07-OCT-1999; 99US-0158029.
PR 08-OCT-1999; 99US-0158232.
PR 12-OCT-1999; 99US-0158369.
PR 13-OCT-1999; 99US-0159293.
PR 13-OCT-1999; 99US-0159294.
PR 14-OCT-1999; 99US-0159329.
PR 14-OCT-1999; 99US-0159330.
PR 14-OCT-1999; 99US-0159331.
PR 14-OCT-1999; 99US-0159637.
PR 14-OCT-1999; 99US-0159638.
PR 18-OCT-1999; 99US-0159584.
PR 21-OCT-1999; 99US-0160741.
PR 21-OCT-1999; 99US-0160767.
PR 21-OCT-1999; 99US-0160768.
PR 21-OCT-1999; 99US-0160770.
PR 21-OCT-1999; 99US-0160814.
PR 21-OCT-1999; 99US-0160815.
PR 22-OCT-1999; 99US-0160980.
PR 22-OCT-1999; 99US-0160981.
PR 22-OCT-1999; 99US-0160989.
PR 25-OCT-1999; 99US-0161404.
PR 25-OCT-1999; 99US-0161405.
PR 25-OCT-1999; 99US-0161406.
PR 26-OCT-1999; 99US-0161359.
PR 26-OCT-1999; 99US-0161360.
PR 26-OCT-1999; 99US-0161361.
PR 28-OCT-1999; 99US-0161920.
PR 28-OCT-1999; 99US-0161992.
PR 28-OCT-1999; 99US-0161993.
PR 29-OCT-1999; 99US-0162142.
```

Query Match 2.28; Score 44; DB 21; Length 746;

Best Local Similarity 51.0%; Pred. No. 0.053;

Matches 104; Conservative 0; Mismatches 100; Indels 0; Gaps 0;

```
QY 542 AGCGCGAGATGACCAACAACTCGGAGAGAGCCCACTCAAGTCAAGATGA 601
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 168 agtggaggaagagagagagagagagagagagagagagagagagagagag 227
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 602 AAACATGAGCAATATGAGCTCTACTACAGACCAAGCCGTTCTGAGGTGAGGAGATGA 661
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 228 aagagagagagagagagagagagagagagagagagagagagagagagag 287
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 662 TTCGAGACATGGTGTGGAGACATCAGCGGTGAGACAGCTGCTGTACTGCGTCC 721
```

```
Db 288 gttacagagagacagtgaggaagaagctgtggagaggtggaagacgcttacttcta 347
QY 722 TCAGCAAGAGACTATGAGCAATCTGA 745
    ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 348 tcaagaatgagcttgatgatctca 371
```

RESULT 11

AA158369 standard; cDNA: 7453 BP.

AA158369;

22-OCT-2001 (first entry)

Human polynucleotide SEQ ID NO 572.

Human; neotropic; immunosuppressant; cytostatic; gene therapy; cancer;

peripheral nervous system; neuropathy; central nervous system; CNS;

Alzheimer's; Parkinson's disease; Huntington's disease; haemostatic;

anyotrophic lateral sclerosis; Shy-Drager Syndrome; chemotactic;

chemokinetic; thrombolytic; drug screening; arthritis; inflammation;

leukaemia; ss.

Homo sapiens.

MO200153312-A1.

26-JUL-2001.

26-DEC-2000; 2000MO-US34263.

21-JAN-2000; 2000US-0488725.

25-APR-2000; 2000US-0552317.

09-JUL-2000; 2000US-0598042.

19-JUL-2000; 2000US-0620312.

03-AUG-2000; 2000US-0653450.

14-SEP-2000; 2000US-0662191.

19-OCT-2000; 2000US-0693036.

29-NOV-2000; 2000US-0727344.

(HYSE-) HYSEQ INC.

Tang YT, Liu C, Asundi V, Chen R, Ma Y, Qian XB, Ren F, Wang D;

Wang J, Wang Z, Wehrman T, Xu C, Xue AJ, Yang Y, Zhang J;

Zhao QA, Zhou P, Goodrich R, Drmanac RT;

WPI: 2001-442253/47.

P-PSDB: AAM39213.

Novel nucleic acids and polypeptides, useful for treating disorders

such as central nervous system injuries -

Claim 1: SEQ ID NO 572; 10078bp; English.

The invention relates to human nucleic acids (AA157798-AA161369) and

the encoded polypeptides (AAM38642-AAM42213) with neotropic,

immunosuppressant and cytostatic activity. The polynucleotides are useful

in gene therapy. A composition containing a polypeptide or polynucleotide

of the invention may be used to treat diseases of the peripheral nervous

system, such as peripheral nervous injuries, peripheral neuropathy and

localised neuropathies and central nervous system diseases, such as

Alzheimer's, Parkinson's disease, Huntington's disease, amyotrophic

lateral sclerosis, and Shy-Drager Syndrome. Other uses include the

utilisation of the activities such as: immune system suppression,

Activin/inhibin activity, chemotactic/chemokinetic activity, haemostatic

and thrombolytic activity, cancer diagnosis and therapy, drug screening,

assays for receptor activity, arthritis and inflammation, leukaemias and

C.N.S disorders.

Note: The sequence data for this patent did not form part of the printed

specification.

DR WPT; 2001-442253/47.

XX

CC	PR	08-NOV-2000;	2000US-0246524.
CC	PR	08-NOV-2000;	2000US-0246525.
CC	PR	08-NOV-2000;	2000US-0246526.
CC	PR	08-NOV-2000;	2000US-0246527.
CC	PR	08-NOV-2000;	2000US-0246528.
CC	PR	08-NOV-2000;	2000US-0246532.
CC	PR	08-NOV-2000;	2000US-0246609.
CC	PR	08-NOV-2000;	2000US-0246610.
CC	PR	08-NOV-2000;	2000US-0246611.
CC	PR	08-NOV-2000;	2000US-0246613.
CC	PR	17-NOV-2000;	2000US-0249207.
CC	PR	17-NOV-2000;	2000US-0249208.
CC	PR	17-NOV-2000;	2000US-0249209.
CC	PR	17-NOV-2000;	2000US-0249210.
CC	PR	17-NOV-2000;	2000US-0249211.
CC	PR	17-NOV-2000;	2000US-0249212.
CC	PR	17-NOV-2000;	2000US-0249213.
CC	PR	17-NOV-2000;	2000US-0249214.
CC	PR	17-NOV-2000;	2000US-0249215.
CC	PR	17-NOV-2000;	2000US-0249216.
CC	PR	17-NOV-2000;	2000US-0249217.
CC	PR	17-NOV-2000;	2000US-0249218.
CC	PR	17-NOV-2000;	2000US-0249244.
CC	PR	17-NOV-2000;	2000US-0249245.
CC	PR	17-NOV-2000;	2000US-0249264.
CC	PR	17-NOV-2000;	2000US-0249265.
CC	PR	17-NOV-2000;	2000US-0249297.
CC	PR	17-NOV-2000;	2000US-0249299.
CC	PR	17-NOV-2000;	2000US-0249300.
CC	PR	01-DEC-2000;	2000US-0250160.
CC	PR	01-DEC-2000;	2000US-0250391.
CC	PR	05-DEC-2000;	2000US-0251030.
CC	PR	05-DEC-2000;	2000US-0251088.
CC	PR	05-DEC-2000;	2000US-0256719.
CC	PR	06-DEC-2000;	2000US-0251479.
CC	PR	08-DEC-2000;	2000US-0251856.
CC	PR	08-DEC-2000;	2000US-0251868.
CC	PR	08-DEC-2000;	2000US-0251869.
CC	PR	08-DEC-2000;	2000US-0251989.
CC	PR	08-DEC-2000;	2000US-0251990.
CC	PR	11-DEC-2000;	2000US-0254097.
CC	PR	05-JAN-2001;	2001US-0259678.
PA	(HUMA-)	HUMAN GENOME SCI INC.	
PT	Rosen CA,	Barash SC,	Ruben SM;
XX	WPI:	2001-483426/52.	
DR	Nucleic acids encoding human immune/hematopoietic antigen polypeptides,		
XX	useful for preventing, diagnosing and/or treating cancers and		
PT	metastasis -		
XX			
XX	Disclosure; SEQ ID NO 36967; 3071bp + Sequence Listing; English.		
XX	AAM54951 to AAK64702 encode the human immune/haematopoietic antigen (I)		
CC	amino acid sequences given in AAM82170 to AAM91921. (I) have cytosolic		
CC	activity, and can be used in gene therapy and vaccine production. (I)		
CC	proteins and polynucleotides may be used in the prevention, diagnosis and		
CC	treatment of diseases associated with inappropriate (I) expression. For		
CC	example, they may be used to treat disorders associated with decreased		
CC	expression by rectifying mutations or deletions in a patient's genome		
CC	that affect the activity of (I) by expressing inactive proteins or to		
CC	supplement the patients own production of (I). Additionally, (I)		
CC	polynucleotides may be used to produce the secreted (I), by inserting		
CC	the nucleic acids into a host cell and culturing the cell to express the		
CC	protein. (I) proteins and polynucleotides may be used to prevent,		
CC	diagnose and treat immune/haematopoietic-related diseases, especially		
CC	cancers and cancer metastases of hematopoietic-derived cells. AAK64703		
CC	to AAK87594 represent human immune/haematopoietic antigen genomic		
CC	sequences from the present invention. AAK54942 to AAK54950 and AAM82169		
CC	represent sequences used in the exemplification of the present invention.		

SQ	Sequence	3229 BP,	830 A;	837 C;	1045 G;	517 T;	0 other:
	Query Match	2.2%;	Score 43.2;	DB 22;	Length 3229;		
	Best Local Similarity	48.4%;	Pred. No.	0.2;	Mismatches 156;	Indels 3;	Gaps 1;
	Matches 151;	Conservative 0;	Mismatches 156;	Indels 3;	Gaps 1;		
OY	544 CGGCAGATGACCAACAACAGCTCGGGAGAGGCCACCAGACTCAAGTGAAGTGAA	603					
Dd	97 caggcggatgagtgcttcgcagctcgggagagctcggaagccactgagcgcgcaagaagc	156					
OY	604 ACCATTGAGCAAAATTGAGCTCCTTA CTCCAGACCAGCCGTTC TGAGTGGAGAAGATGATT	663					
Dd	157 gcccgaggagcgacgaggtcctcctcctcgcagcagcgtgtgcccgccttgcgtgaagaaag	216					
OY	664 CCAGACATGCGGTGGGAGCACGACCGGCGAGCGACGCTGCTGTGATGCGCTGTCCC---	720					
Dd	217 agcgcggcgccaggaagaagttcacagaagaagagtgtgtgaacctcagaatagcccccag	276					
OY	721 CTCAGAAAAGACTATGAGAAATCTGAAGGAAGCTCGSAAAGCCACAGGGGAACCTGCTGAC	780					
Dd	277 ctggagcgcaagattaccagcagctgcagagagacccacccagcgccagagaccagctcaggag	336					
OY	781 AGCTGAAGAAGGATTTGCTGCTCTAGGACCAAGTTGACACTCTCAACACTGAGCTG	840					
Dd	337 aagcaggaggaagagctgactcctccagacaagctccaagaagctagaagaagagagcgg	396					
OY	841 GATCAGGCCAAG	852					
Dd	397 gccatgccccgag	408					
	RESULT 16						
	AAS69541						
ID	AAS69541 standard; CDNA: 390 BP.						
XX	AAS69541:						
XX	13-FEB-2002 (first entry)						
DE	DNA encoding novel human diagnostic protein #5345.						
XX	Human; chromosome mapping; gene mapping; gene therapy; forensic;						
KW	Food supplement; medical imaging; diagnostic; genetic disorder; ss.						
OS	Homo sapiens.						
XX	WO200175067-A2.						
FN	11-OCT-2001.						
PD	30-MAR-2001; 2001WO-US08631.						
PF	31-MAR-2000; 2000US-0540217.						
PR	23-AUG-2000; 2000US-0649167.						
XX	(HXSE-) HXSEQ INC.						
PA							
XX							
PI	Dzmanec RT, Liu C, Tang YT;						
XX							
DR	WI: 2001-639362/73.						
XX	P-PSDB: ABG05354.						
PT	New isolated polynucleotide and encoded polypeptides, useful in						
PT	diagnostics, forensics, gene mapping, identification of mutations						
PT	responsible for genetic disorders or other traits and to assess						
PT	biodiversity -						
PS	Claim 1; SEQ ID NO 5345; 103pp: English.						
XX							
CC	The invention relates to isolated polynucleotide (I) and						
CC	polypeptide (II) sequences. (I) is useful as hybridisation probes,						
CC	polymerase chain reaction (PCR) primers, oligomers, and for chromosome						

CC and gene mapping, and in recombinant production of (II). The
CC polynucleotides are also used in diagnostics as expressed sequence tags
CC for identifying expressed genes. (I) is useful in gene therapy techniques
CC to restore normal activity of (II) or to treat disease states involving
CC (II). (II) is useful for generating antibodies against it, detecting or
CC quantitating a polypeptide in tissue, as molecular weight markers and as
CC a food supplement. (II) and its binding partners are useful in medical
CC imaging of sites expressing (II). (I) and (II) are useful for treating
CC disorders involving aberrant protein expression or biological activity.
CC The polypeptide and polynucleotide sequences have applications in
CC diagnostics, forensics, gene mapping, identification of mutations
CC responsible for genetic disorders or other traits to assess biodiversity
CC and to produce other types of data and products dependent on DNA and
CC amino acid sequences. AAS64197-AAS94564 represent novel human
CC diagnostic coding sequences of the invention.
CC Note: The sequence data for this patent did not appear in the printed
CC specification, but was obtained in electronic format directly from WIPO
CC at http://wipo.int/pub/published_pct_sequences.
SQ Sequence 390 BP; 191 A; 14 C; 149 G; 36 T; 0 other;

Query Match 2.2%; Score 42.8; DB 23; Length 390;
Best Local Similarity 46.6%; Pred. No. 0.081;
Matches 137; Conservative 0; Mismatches 157; Indels 0; Gaps 0;

QY 629 TCCAGAGCCAGCTCTGAGGAGATGATTCGAGCATGGGCTGGACAGTCAG 688
Db 33 tccaacacagagcagcatcagctcggaggaaggaaggaaggaaggaaggaag 92
QY 689 CGGTGGAGCAGCTGCTGTCTACTGCGTGTCCCTCAAGAAAGATGTGAAATCTGNAAG 748
Db 93 aggaaggaaggaaggaaggaaggaaggaaggaaggaaggaaggaaggaag 152
QY 749 AACCTGGAAAGCCACAGGGGAAGCTGCTGACAGGTTGAAGAGATTGCTCTCTA 808
Db 153 aggaaggaaggaaggaaggaaggaaggaaggaaggaaggaaggaagga 212
QY 809 GAGGACAGTTGAGACTCTCAACACAGCTGATGATCAGGCCAGTTAGACTAGCTGAC 868
Db 213 ageggaaggaaggaaggaaggaaggaaggaaggaaggaaggaaggaag 272
QY 869 CCCAGAGGACTTACAAAGTGTGACAGGAGATCAGACCTTACGAAAGTAAT 922
Db 273 aggaaggaaggaaggaaggaaggaaggaaggaaggaaggaaggaag 326

RESULT 17

AAT05868
ID AAT05868 standard; DNA; 3399 BP.

XX AAT05868;
XX
XX 14-AUG-1996 (first entry)
XX
XX Chicken leucocytozoan DNA encoding immunogenic protein for vaccines.
DE
XX
XX Chicken leucocytozoan; immunogen; recombinant vaccine; protection;
KW Immunisation; vaccination; ss.
XX
XX Chicken leucocytozoan.

XX Key Location/Qualifiers

XX CDS 1..3399

XX FT misc_feature 1150..33218

XX FT /tag= a

XX FT /tag= b

XX FT /note= "Fragment referred to in the claims, for
XX use as insert in a recombinant vaccine
XX against chicken leucocytozoan disease"

XX JP07284392-A.

PD 31-OCT-1995.
XX
XX 19-APR-1994; 94JP-0080643.
XX
XX 19-APR-1994; 94JP-0080643.
XX
XX (DOHU-) DOBUTSUYO SEIBUTSUGAKUTEKI SEIZAI KYOKAI.
PA (KITA) KITASATO KENKYUSHO SH.
XX
XX WPI: 1996-006311/01.
DR P-PSDB: AAR97866.
DR
XX
XX Chicken leucocytozoan immunogenic protein - used in a recombinant
PT vaccine against chicken leucocytozoan disease
XX
PS Claim 6; Page 6-9; 35pp; Japanese.
XX
XX AAT05868 encodes a chicken leucocytozoan immunogenic protein, this DNA
CC or a fragment of it can be used in a recombinant vaccine to immunise
CC against chicken leucocytozoan disease. The DNA is used in a vector
CC and operatively linked to an expression regulatory sequence as in
CC standard practice.
SQ Sequence 3399 BP; 1577 A; 508 C; 798 G; 516 T; 0 other;

Query Match 2.2%; Score 42.8; DB 17; Length 3399;
Best Local Similarity 43.5%; Pred. No. 0.26;
Matches 194; Conservative 0; Mismatches 252; Indels 0; Gaps 0;

QY 476 AGAACCCCTTAAACAGGACGAGATGCTGTGTTCCACCCCTGAAAAACAGATGAATGCC 535
Db 2349 agtaacacatgaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2408
QY 536 TGGAGCAGCCGACGATGAGACCAACAAAGCTCGGAGAGAGCCACCGACTTCAGTGCA 595
Db 2409 acatgaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2468
QY 596 AGATGAAACCATGAGCAATTTGACTCTCTACTCCAGACCCGTTCTGAGGTGAGG 655
Db 2469 agaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaa 2528
QY 656 AGATGATTCGAGATGAGGCTGTGAGACAGTCAGCGTGGACAGCTGCTGACTGCG 715
Db 2529 aaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2588
QY 716 TGTCCCTCAAGAAAGATGAGATCTGAAAGAACTCGGAAGGCCACAGGGAACTGG 775
Db 2589 tgaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2648
QY 776 CTGACAGTTGAAGAGATTTGCTGCTCTTAGGACCAAGTTGAAGACTCTCAACACTG 835
Db 2649 tgaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2708
QY 836 AGCTGATCAGGCCAAGTTGAACTGAGCTGAGTCACCCAGAAAGACTTACAAAGTGTGACC 895
Db 2709 aaaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaagaaag 2768
QY 896 AGGAGATCAGGACCTTAAAGAAAGAG 921
Db 2769 agaaagaaagaaagaaagaaagaaag 2794

RESULT 18

AB199537
ID AB199537 standard; cDNA; 1080 BP.

XX AB199537;

XX 07-MAR-2002 (first entry)

XX Mouse ischaemic condition related cDNA sequence SEQ ID NO:537.

KM Mouse; ischaemia; compressive ischaemia; occlusive ischaemia;
 KM vasospastic ischaemia; ischaemic condition; ischaemic disease; ss.
 OS Mus musculus.
 PN W0200188188-A2.
 PD 22-NOV-2001.
 PF 18-MAY-2001; 2001MO-JP04192.
 PR 18-MAY-2000; 2000JP-0145977.
 PA (UNIV-) UNIV NIHON SCHOOL JURIDICAL PERSON.
 XX Ishikawa K, Asai S, Takahashi Y, Nagata T, Ishii Y;
 DR WPI; 2002-034733/04.
 DR P-PSDB; ABB57221.
 XX
 PT Examining the ischemic condition (e.g. occlusive ischemia) by measuring
 PT expression levels of particular genes defined in the specification or
 PT by determining the expression profile of a gene group comprising these
 PT genes -
 PS
 PS Claim 2; Page 1472-1473; 2690pp; English.
 CC The present invention describes a method for examining ischaemic
 CC conditions, comprising measuring the expression levels of particular
 CC genes (1) in a test sample or determining the expression profile of a
 CC gene group in the sample comprising genes selected from (1). The method
 CC is useful for examining the ischaemic condition (e.g. compressive
 CC ischaemia, occlusive ischaemia or vasospastic ischaemia) by measuring
 CC expression levels of particular genes (AB199202 to AB199912, encoding
 CC the protein sequences in ABB57020 to ABB57374) or by determining the
 CC expression profile of a gene group comprising these genes. The
 CC expression levels or expression profiles produced by these genes are
 CC used as an indicator when screening for ischaemic condition-improving
 CC drugs or therapeutics for ischaemic diseases. AB199913 and AB199914
 CC represent PCR primers for a mouse ischaemic condition related sequence,
 CC which are used in the exemplification of the present invention.
 CC
 XX
 XX Sequence 1080 BP; 370 A; 191 C; 390 G; 129 T; 0 other;
 SQ
 Query Match 2.1%; Score 42.4; DB 24; Length 1080;
 Best Local Similarity 45.3%; Pred. No. 0.18;
 Matches 134; Conservative 0; Mismatches 186; Indels 0; Gaps 0;
 QY 517 AAAAAGCAGATGAAGTTCCTGAGACGCGGAGATGAGACCAAAAGCTCGGAGAG 576
 |||| | | | | |||| | | | | |||| | | | | |||| | | | |
 Db 703 aaaaagaagaagaagaagaagaagaagaagaagaagaagaagaagaagaaga 762
 QY 577 GCCCACCAGCTCAAGTGCAGATGAAAACCATGAGCAAAATTGACTCTACACAGC 636
 | | | | | | | | | | | | | | | | | | | | | | | |
 Db 763 gagagaagaagaagaagaagaagaagaagaagaagaagaagaagaagaag 822
 QY 637 CACCGTTCTAGAGTGGAGGAGATGATTCGAGACATGGGTGTGGACACGTACGCGTGGAG 636
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 823 aagagaagaagaagaagaagaagaagaagaagaagaagaagaagaagaag 882
 QY 697 CACCTGCGCTTGTACTCCGTCCTCCATCAGAAAGATATGAGAACTGAGAGCTCGG 756
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 883 aagagaagaagaagaagaagaagaagaagaagaagaagaagaagaagaag 942
 QY 757 AAGGCCACAGGGAACTGCTGACAGGTTGAAGAAGGATTTGGTCTCTAGAGCAAG 816
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 943 aagagaagaagaagaagaagaagaagaagaagaagaagaagaagaagaag 1002
 QY 817 TTGAAGACTCTCAACACTGAGCTGGATCAAGCCAACTTAG 856
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 1003 aagagaagaagaagaagaagaagaagaagaagaagaagaagaagaag 1042

RESULT 19
 ID AAN71065 standard; DNA: 2000 BP.
 XX
 AC AAN71065;
 XX
 DT 01-JAN-1980 (first entry)
 XX
 DE Gene encoding Plasmodium cynomolgi sporozoite circumsporozoite
 DE protein.
 XX
 KM Immunogen; vaccine; malaria; immunodominant epitope; DNA probe; ss.
 KM
 OS Plasmodium cynomolgi.
 XX
 FH Location/Qualifiers
 FT CDS 326..1534
 FT /tag- a
 FT /product- circumsporozoite protein
 FT
 PN W08700533-A.
 XX
 PD 29-JAN-1987.
 XX
 PF 24-JUN-1986; 86MO-US01373.
 XX
 PR 12-JUL-1985; 85US-0754645.
 XX
 PA (UNY-) NEW YORK UNIV.
 PA (ARNO/) ARNOT D E.
 XX
 PI Arnot DE, Enea V, Nussenzwei RS, Nussenzweig V;
 XX
 DR WPI; 1987-037250/05.
 DR P-PSDB; AAP70709.
 XX
 PT New Plasmodium vivax circumsporozoite protein - and synthetic
 PT peptide(s) contg. its dominant epitope, useful in anti-malarial
 PT vaccines
 PT
 PS Disclosure; fig. 6; 32pp; English.
 XX
 CC The gene encoding the circumsporozoite protein of P. cynomolgi
 CC is used during the detection of the circumsporozoite protein of P.
 CC vivax. A DNA probe from this P. cynomolgi sequence was a 700 bp
 CC PstI fragment encoding the C-terminal domain and about 350
 CC bases of the 3' untranslated region of the gene. Specifically,
 CC the probe, designated P236-7, encompasses bases 851-1827 flanked
 CC by C and G tails. This gene is useful in the construction of an
 CC anti-malarial vaccine. See also AAP70704-08 and AAN71064.
 CC
 XX
 XX Sequence 2000 BP; 655 A; 318 C; 527 G; 500 T; 0 other;
 SQ
 Query Match 2.1%; Score 41.6; DB 8; Length 2000;
 Best Local Similarity 43.5%; Pred. No. 0.43;
 Matches 188; Conservative 0; Mismatches 244; Indels 0; Gaps 0;
 QY 370 AAGCTCAGCTTCCCGAAGACAGGAGAAACGGAGACGCCGCTTATCGACT 429
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 531 aaggaagctgataaaccaaaaaaagaagcaaaaaaagtagaacaacaaaaaagcaag 590
 QY 430 CTACGGGACACCTTGAAGAACCAATGCTACCGTGGAGTCCCTACAGAACCTTAAAC 489
 | | | | | | | | | | | | | | | | | | | | | |
 Db 591 cgaaaaaataagctgaaacaacaaagaagatgagctgctgcaagaagaagaaga 650
 QY 490 AAGCAGAGATGCTGTGTTCCACCTGGAAGAAACAGATGAAGTCTGAGCAGCGGAG 549
 || | | | | | | | | | | | | | | | | | | | | | |
 Db 651 atgattgagctgctgctgcaagaagaagaagaagaatgattgagctgctgcaag 710
 QY 550 GATGAGACCAAAACACCTCGGAGGAGGCCACCGACTCAAGTGAAGTGAAGAACCAAG 609
 || | | | | | | | | | | | | | | | | | | | | | |

[illegible]

DT	23-FEB-2001	(first entry)
XX	Human I-kappa-B kinase gamma-subunit (IKK-gamma) cDNA.	
XX	Human: I-kappa-B kinase; IKK; antisense therapy; gene therapy;	
XX	cytokine expression inhibition; NF-kappa-B activation inhibition;	
KW	nuclear factor-kappa-B; rheumatoid arthritis; immune disorder;	
KW	cancer; IKK-gamma; gamma-subunit; ss.	
OS	Homo sapiens.	
XX	JP2000253884-A.	
PN	19-SEP-2000.	
PD	10-MAR-1999; 99JP-0063291.	
XX	10-MAR-1999; 99JP-0063291.	
XX	(TOAG) TOA GOSEI CHEM IND LTD.	
PA	WPI: 2000-655813/64.	
DR	Antisense nucleic acid compound complementary to the subunit of	
XX	Ikappab, used to treat rheumatic arthritis, immune diseases and cancer	
PT	-	
XX	Claim 3; Page 14-15; 20pp; Japanese.	
PS	The invention relates to an antisense oligonucleotide targeted to	
XX	a gene encoding a subunit of I-kappa-B kinase (IKK) which inhibits its	
CC	expression, and thereby inhibits expression of a cytokine such as	
CC	IL-6 (interleukin-6). I-kappa-B kinase activates NF-kappa-B (nuclear	
CC	factor-kappa-B) which acts a transcriptional regulator of cytokine	
CC	genes. The antisense oligonucleotide can be used in gene therapy to	
CC	treat rheumatoid arthritis, immune disorders and cancers. Sequences	
CC	AA681422-C81426 are cDNAs derived from genes whose expression may be	
CC	inhibited using an antisense oligonucleotide of the invention.	
CC	The present sequence represents a human IKK-gamma subunit cDNA.	
XX	Sequence 1994 BP; 429 A; 585 C; 633 G; 347 T; 0 other;	
SO		
Query Match	2.1%; Score 40.8; DB 21; Length 1994;	
Best Local Similarity	46.2%; Pred. No. 0.73;	
Matches 135; Conservative 0; Mismatches 157; Indels 0; Gaps 0;		
OY	528 GAAGTTCCTGGAGCGGAGCATGATGACCAAGACGCTGGAGGAGGCCACGACT 587	
DB	961 ggaagcccccggagcccaacagaggtgctgctagctgaagtgagagccgagcaaga 1020	
OY	588 CAAGTGAAGATGAATAACCATGAGACAAATTGAGCTCTACTCCAGAGCCAGCTTCTGA 647	
DB	1021 caagatctgtatgtagagaccgttcgcgtgctgaagagcccgagtgatctacaagcgaga 1080	
OY	648 GGTGAGGAGATGATTTCCAGACATGGCTGTGGACACAGTCACGCGGTGGAGCAGCTGGCTGT 707	
DB	1081 ctccagagctgagagagccagccgggagaaagcttgcagagaaagagagagctctgcagga 1140	
OY	708 GTACTGCTGTGCCCTCAGAAAGATATGACAAATGTGAAGAAAGCTCGAAGGCCACAGG 767	
DB	1141 gcaagcttgagacagctgcgcagagagatcacagcaactgaagagcagctgctcaggaatcgagc 1200	
OY	768 GGAACCTGGCTGACAGGTTGAAGAAAGATTTGGTGTGCTCTAGACAGCAAGTTG 819	
DB	1201 cagagctcgagagcatgtagaagcagcagctgctcagagctcccgagcccttg 1252	
RESULT	23	
AAA35027		
ID	AAA35027 standard; DNA; 1994 BP.	
NC	AAA35027;	

XX	28-JUL-2000 (first entry)
DE	
XX	Human adenosine receptor related polynucleotide SEQ ID NO:2716.
XX	
KW	Human; adenosine receptor; low adenosine antisense oligonucleotide;
KW	phosphorothioate; impaired respiration; inflammation; allergy;
KW	allergic diseases; bronchoconstriction; inhibitor; antiinflammatory;
KW	antiallergic; antiasthmatic; cyostatic; analgesic; impaired airway;
KW	lung disease; ischemic condition; pulmonary vasoconstriction; asthma;
KW	respiratory distress syndrome; pain; cystic fibrosis; emphysema;
KW	pulmonary hypertension; chronic obstructive pulmonary disease; COPD;
KW	cancer; leukaemia; lymphoma; carcinoma; metastasis; ss.
XX	
OS	Homo sapiens.
XX	
PN	WO200009525-A2.
XX	
PD	24-FEB-2000.
XX	
PF	03-AUG-1999; 99WO-US17712.
XX	
PR	03-AUG-1998; 98US-0095212.
XX	
PA	(UYEC-) UNIV EAST CAROLINA.
XX	
PI	Nyce JW;
DR	WPI: 2000-205971/18.
XX	
PT	New antisense oligonucleotides useful for treating e.g. pulmonary
PT	vasoconstriction, inflammation, allergies, asthma, hypertension,
PT	bronchitis, emphysema, respiratory distress syndrome, ischemia or
XX	cancers -
XX	
PS	Disclosure; Page 968-969; 1343pp: English.
XX	
CC	The present invention describes a new composition comprising an
CC	antisense oligonucleotide (ON) with low adenosine (up to 15%), which
CC	targets nucleic acids involved in bronchoconstriction, allergies, and/or
CC	inflammation. The ON can have antiinflammatory, antiallergic,
CC	antiasthmatic, cytostatic and analgesic activities. The compositions are
CC	useful for the treatment of diseases associated with inflammation,
CC	impaired airways, including lung disease and diseases whose secondary
CC	effects afflict the lungs of a subject. They can be used for treating
CC	e.g. ischemic conditions, pulmonary vasoconstriction, allergies,
CC	asthma, impeded respiration, respiratory distress syndrome, pain, cystic
CC	fibrosis, pulmonary hypertension, emphysema, chronic obstructive
CC	pulmonary disease (COPD), and cancers such as leukemias, lymphomas,
CC	carcinomas, and cancers which may metastasize to the lungs, including
CC	breast and prostate cancer. The reduction of the adenosine content of
CC	the ONS reduces side effects. The A-containing ONS break down with the
CC	release of deoxyadenosine which activates adenosine receptors causing
CC	bronchoconstriction and inflammation. AAA32313 to AAA3312 represent the
CC	nucleotide sequences given in the sequence listing from the present
CC	invention, which correspond to SEQ ID NO:1 to 2815, and then the last
CC	185 sequences are also called SEQ ID NO:1 to 185, and the sequences
CC	differ from the previously named sequences. SEQ ID NO:11 to 1680
CC	(AAA33223 to AAA33992) are specifically claimed ONS from the present
CC	invention. N.B. Sequences given in the disclosure of the present
CC	invention do not match up with their corresponding SEQ ID NO: sequences
CC	given in the sequence listing.
XX	
SO	Sequence 1994 BP: 429 A; 585 C; 633 G; 347 T; 0 other;
XX	
QY	Query Match 2.1%; Score 40.8; DB 21; Length 1994; Best Local Similarity 46.2%; Pred. No. 0.73; Matches 135; Conservative 0; Mismatches 157; Indels 0; Gaps
DB	528 GAAGTTCCTGGAGCGACGCGCATGTAGATCAAAACAAGCTCGGAGAGAGGCCACCAGACT 567 961 ggaagccctcgttggccaacagcgatgcataaaagtcaagaaggccagacgaca 1020

YY 588 CAACTGCAAGTCGAGAAAACCATGTAGCAAAATPAGGCTCCACTCCAGACCCAGCGTTCGA 647
 Db 1021 caagattctgctgaggaagccgcttcctcggctctcgaagcccgagcgtatcctcaagcgga 1080
 QY 648 GGTGAGAGCAATATTCATTCGACACATGCGTGTGGACAGTCAGCGTGGAGCAGCTGCGTGT 707
 Db 1081 ctccagagctcgagagcagcgccggagagctcggccggaagaagagagctccctcgagga 1140
 QY 708 GTACTGCGTGTCCCTCAGAGAAGATATAGATCTGGAAGAGCTCGGAAGCCACAGC 767
 Db 1141 gcaagcttgagcaagcagctgcagagaggtacagcaaacctgaagcgagctgtcagagatcgcc 1200
 QY 768 GGAAGCTGGCTGCAGCAGGTGTGAAGAAGATTTGGTGTCTTAGACACAGTTG 819
 Db 1201 cagagatcgagagacatgagagaagcgagcatgtcagatctccagagccctctg 1252

RESULT 24
 AAZ07513
 ID AAZ07513 standard; DNA; 2009 BP.
 AC AAZ07513;
 DT 26-NOV-1999 (first entry)
 XX
 DE Human RIP-associated protein (RAP-2) encoding DNA.
 XX
 KW Receptor interacting protein; RIP-associated protein-2; RAP-2; RIP;
 KW Inflammation; cell death; cell survival; septic shock; hepatitis;
 KW graft versus host rejection; diabetes; multiple sclerosis; tumor;
 KW HIV infection; p55-receptor; FAS-receptor; human; ss.
 XX
 OS Homo sapiens.
 XX
 PN W09947672-A1.
 PD 23-SEP-1999.
 XX
 PF 18-MAR-1999; 99WO-IL00158.
 PR 19-MAR-1998; 98IL-0123758.
 PR 01-SEP-1998; 98IL-0126024.
 XX
 PA (YEDA) YEDA RES & DEV CO LTD.
 PA (YESH) UNIV YESHIVA EINSTEIN COLLEGE.
 XX
 PI Wallach D, Kovalenko A, Horwitz MS, Li Y;
 DR WPI: 1999-562113/47.
 P-PSDB: AAY27430.
 XX
 PT New receptor interacting protein-associated protein-2, used to develop
 PT products for treating, e.g. septic shock, tumors or HIV infection -
 XX
 PS Claim 4; Fig 1A-B; 132pp; English.
 CC
 XX This DNA encodes a receptor interacting protein (RIP)-associated protein
 CC -2 (RAP-2). The RAP-2 proteins, isoforms, analogs, fragments or
 CC derivatives or DNA can be used for the modulation or mediation of the
 CC RIP modulated/mediated intracellular effects on the inflammation, cell
 CC death or cell survival pathways in which RIP is involved directly, or
 CC indirectly via other modulators/mediators of these pathways. They can be
 CC used for treating e.g. septic shock, graft versus host rejection, acute
 CC hepatitis, diabetes or multiple sclerosis. They can also be used for
 CC treating tumor cells or HIV-infected cells or other diseased cells. The
 CC RAP-2 binding proteins can also be used for modulating/mediating the function
 CC of RAP-2. The products can also be used for diagnostic purposes, e.g. for
 CC identifying disorders related to abnormal functioning of cellular effects
 CC mediated by the p55-R, FAS-R or other related receptors.
 XX
 SO Sequence 2009 BP; 418 A; 587 C; 643 G; 356 T; 5 other;

[illegible][illegible]

```
QY 568 CAAGTGAAGATGAAAACCATGAGCAAAATTGAGCTCTACTCCAGAGCCACGCTTTCGA 647
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 7658 caagctgtgagtggagccgcttcgctgtaaggccagcgagatctatcaagaagcgga 7717
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 648 GGTGGAGAGATGATTCCAGACATGGGTGTGGACAGTGCAGGTGGAGCAGCTGGCTGT 707
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 7718 ctccagagctgagagagccgaggaagctggccgagagaagagagctctctcagga 7777
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 708 GTACTGCGTGTCCCTCAGAGAAAGATGAGAAATCTGAAGAGAGCTCGAAGGCCACAGG 767
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 7778 gcaagctgagagcagctgcagagaggaagtaacagcaactgaagccagctgtcaggaagtcg 7837
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 768 GCAACTGGCTGACAGCTGGAAGAGATTGGTGTCTCTAAGAGCAAGTTG 819
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 7838 cagagtcgagagcagatgaggaagcgagctgtcagagtcctccagagcccttg 7889
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||

RESULT 28
AAA30290
ID AAA30290 standard; DNA; 3489 BP.
XX
AC AAA30290:
XX
DE 11-SEP-2000 (first entry)
XX
XX Kaposi's sarcoma-associated herpesvirus LANA gene.
XX
KW Kaposi's sarcoma-associated herpesvirus; KSHV; rhadino virus;
KW latency-associated nuclear antigen; LANA; gamma-2 herpes virus;
KW Human herpes virus 8; HHV8; rhadino virus cis-acting element; RVCAE;
KW Kaposi's sarcoma; primary effusion lymphoma; PEL;
KW human immunodeficiency virus; HIV; multicentric Castleman's disease; ds.
XX
OS Kaposi's sarcoma-associated herpesvirus.
XX
FH Key Location/Qualifiers
FT CDS 1..3489
FT /tag- a
FT /product= "LANA"
FT /tag- b
FT /note= "nuclear localisation signal, NLS"
FT /tag- c
FT /note= "nuclear localisation signal, NLS"
FT misc_signal 190..210
FT misc_signal 190..210
FT /tag- c
FT /note= "nuclear localisation signal, NLS"
XX
PN WO200029626-A1.
XX
PD 25-MAY-2000.
XX
PE 19-NOV-1999; 99WO-US27508.
XX
PR 19-NOV-1998; 98US-0109422.
PR 21-APR-1999; 99US-0298568.
XX
PA (KIEF/) KIEFF E. D.
PA (BALI/) BALLESTAS M E.
PA (KAYE/) KAYE K M.
XX
PI Kieff ED, Ballestas ME, Kaye KM;
XX
XX WPI: 2000-387829/33.
XX DR P-PSDB: AAY96255.
XX
XX Treating or preventing a disease associated with rhadino virus
XX infection in a mammal which includes Kaposi's Sarcoma and Primary
XX Effusion Lymphoma
XX
XX Disclosure: Fig 6; 70pp; English.
XX
XX The present sequence is the Kaposi's sarcoma-associated herpesvirus,
XX (KSHV) latency-associated nuclear antigen (LANA) gene. KSHV is also known
XX as Human Herpes Virus 8 (HHV8) and belongs to the rhadino virus, or
XX
```

```
CC gamma-2 herpes virus class. The LANA protein is necessary for the
CC efficient persistence of rhadino virus DNA in mammalian cells. Persistent
CC rhadino virus infection is implicated in a variety of diseases e.g.
CC Kaposi's Sarcoma (KS), Primary Effusion Lymphoma (PEL) and multicentric
CC Castleman's disease. In addition, KS is a common malignancy in HIV
CC patients. KSHV persists in host cells in a latent form. One of the few
CC genes expressed from the latent viral DNA is LANA. LANA associates with
CC both human chromosomes and with the rhadino virus cis-acting element
CC (RVCAE), thereby providing a tethering function: the KSHV DNA episome is
CC "tied" to the host chromosomes. This allows the viral DNA to persist in
CC the host cell. The present sequence may be used to screen and identify
CC molecules that inhibit LANA interaction with RVCAE, thereby interfering
CC with the latency cycle of this virus. Potential antiviral treatments for
CC the above mentioned diseases may therefore be based on LANA deregulation.
XX
XX Sequence 3489 BP; 1053 A; 862 C; 1137 G; 437 T; 0 other;
XX

Query Match 2.0%; Score 40.4; DB 21; Length 3489;
Best Local Similarity 44.0%; Pred. No. 1.3;
Matches 170; Conservative 0; Mismatches 216; Indels 0; Gaps 0;

QY 514 CTGAAAAAAGATGAGATCTCTGGAGAGCGGAGATGAGACCAAAAGCTCGGAG 573
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2191 cagcagagatgagcagcagcagcagatgagcagcagcagcagcagcagcagcag 2250
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 574 GAGGCCACCGACCTCAAGATGCAAGATGCAAAACATGAGCAAAATTGAGCTCTACTCCAG 633
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2251 gatbaacagagagcagcagagagagagcagcagcagcagcagcagcagcagcagcag 2310
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 634 AGCCAGCTGTGAGCTGGAGAGATGATTCGAGCATGGCTGTGGACAGTCAGCGGTG 693
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2311 gagcagagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagc 2370
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 694 GAGCAGCTGCTGTGTACTGTGCTCTCCCTCAAGAAAGATGTGAGATCTAAGAGAGCT 753
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2371 gagagagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagc 2430
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 754 CGGAAGCCACAGGAGCACTGCTGACAGTGTGAAGAGATTTGCTCTCTAGAGAGC 813
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2431 tttagagagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagc 2490
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 814 AAGTTGAAGACTCTCAACACTGAGCTGATCAGGCCAAGTTAGAACTGAGTCAAGCCAG 873
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2491 gacttagagagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcag 2550
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
QY 874 AAGCACTTACAAAGTGTGACCAGCA 899
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||
Db 2551 gtggaagagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagc 2576
      ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| ||||| |||||

RESULT 29
AAF82901
ID AAF82901 standard; DNA; 3489 BP.
XX
AC AAF82901:
XX
DE 29-JUN-2001 (first entry)
XX
XX Nucleotide sequence of KSHV tethering protein, LANA.
XX
XX Histone H1; tethering protein; LANA; gene therapy; multiple sclerosis;
XX Parkinson's disease; Huntington disease; diabetes; human herpesvirus 8;
XX KSHV; latency-associated nuclear antigen; LANA; ds.
XX
XX Kaposi's sarcoma associated herpesvirus.
XX
XX Key Location/Qualifiers
XX CDS 1..3489
XX /tag- a
XX
XX WO200125484-A2.
XX
```



```

Oy 754 CGAAGCCACAGGGGACTGCTGCTACAGTTCAGAGAGATTGGTCTCTAGAGAC 813
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 19566 TTAGAGACAGCAGACAGAGATTAGAGCAGAGAGATTAGAGAGAGAGAGCAG 19507
Oy 814 AAGTTGAAGACTCTCAACACTGACTGCATCGAGCCAGTTGAACTGAGTCCAGCCAG 873
    |||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 19506 GAGTTAGAGAGCAGCAGAGAGATTAGAGAGAGAGAGAGAGAGAGAGAGAG 19447
Oy 874 AAGGACTTACAAAGTGTGACCAGCA 899
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 19446 GTGGAAGAGCAAGACAGAGAGAGCA 19421

RESULT 31
AAV19941/C
ID AAV19941 standard; DNA; 137507 BP.
XX
XX AAV19941;
AC
XX
XX 03-AUG-1998 (first entry)
DT
XX
XX KSHV Long unique coding region and terminal repeat.
DE
XX
XX KSHV; HHV8; human herpes virus 8; macrophage inflammatory protein II;
KW Interleukin-6; IL-6; interferon regulatory factor; rheumatoid arthritis;
KW complement-binding protein; glycoprotein; capsid protein IV; infection;
KW immediate early protein; Kaposi's sarcoma; protective vaccine; lymphoma;
KW lymphoproliferative disease; leukaemia; splenomegaly; mycosis fungoides;
KW HIV immune status; anti-inflammatory agent; therapy; ds.
XX
XX Kaposi's sarcoma-associated herpes virus.
OS
XX
XX Key Location/Qualifiers
FH 1142..2794
FT /*tag= a
FT /product= complement-binding protein
FT 8699..11236
FT /*tag= b
FT /product= glycoprotein B
FT complement (17261..17875)
FT /*tag= c
FT /product= interleukin 6
FT complement (21548..21832)
FT /*tag= d
FT /product= macrophage inflammatory protein II
FT complement (27137..27424)
FT /*tag= e
FT /product= interferon regulatory factor 1
FT 28661..29741
FT /*tag= f
FT /product= protein TI.1
FT complement (58976..60175)
FT /*tag= g
FT /product= glycoprotein M
FT complement (69412..69915)
FT /*tag= h
FT /product= glycoprotein L
FT complement (88410..88910)
FT /*tag= i
FT /product= interferon regulatory factor 2
FT 89600..90541
FT /*tag= j
FT /product= interferon regulatory factor 3
FT 90173..90643
FT /*tag= k
FT /product= glycoprotein X
FT complement (93636..94127)
FT /*tag= l
FT /product= interferon regulatory factor 4
FT complement (111931..112443)
FT /*tag= m
FT /product= capsid protein IV
FT complement (123808..127296)
FT
FT CDS

```

```

FT /*tag= n
FT /product= immediate early protein
FT
FM W09804576-A1.
XX
XX 05-FEB-1998.
PD
XX
PF 22-JUL-1997; 97WO-US13346.
XX
PR 29-NOV-1996; 96US-0757669.
PR 25-JUL-1996; 96US-0686243.
PR 25-JUL-1996; 96US-0686349.
PR 25-JUL-1996; 96US-0686350.
PR 25-JUL-1996; 96US-0687253.
PR 25-JUL-1996; 96US-0688814.
PR 05-SEP-1996; 96US-0708678.
PR 10-OCT-1996; 96US-0728323.
PR 13-NOV-1996; 96US-0747887.
PR 13-NOV-1996; 96US-0748640.
XX
XX (UYCO ) UNIV COLUMBIA NEW YORK.
PA
XX
PI Bohenzky RA, Chang Y, Edelman IS, Moore PS, Russo JJ;
XX
XX WPI; 1998-130615/12.
DR
XX
XX New nucleic acid encoding Kaposi's sarcoma associated herpes virus
PT proteins - useful for, e.g. detecting levels of HHV8 in, and
PT preparation of vaccines for treatment of, HIV patients
PT
XX
XX Example 2; Page 135-203; 230pp; English.
PS
XX

```

This sequence represents the long unique region and terminal repeat of the Kaposi's sarcoma-associated herpes virus (KSHV). KSHV is also known as human herpes virus 8 (HHV8). This sequence contains the DNAs of the invention which encode KSHV polypeptides selected from: (a) viral macrophage inflammatory protein (MIP) II; (b) viral interleukin-6 (IL-6); (c) viral IRF 1; (d) complement-binding protein; glycoproteins B, M or L; (d) capsid protein IV encoded by ORF65; and (e) immediate early protein encoded by ORF73. Labelled probes for the nucleic acid, proteins encoded by it, and antibodies (Ab) specific for the proteins are useful for detecting HHV8, specifically for diagnosis of Kaposi's sarcoma, in body fluids or tissue samples. HHV8 infections can be treated with antisense or triplex forming molecules or agents that bind specifically to the protein. Ab may be used for prophylaxis or treatment of HHV8 infection, while the protein can be used in protective vaccines. Ab may also be used to differentiate between lymphomas, and HHV8 may be implicated in many other lymphoproliferative diseases such as lymphomas, leukaemia, splenomegaly and mycosis fungoides. Cells and animals containing the nucleic acid are useful for drug screening. HHV8-derived nucleic acid can be used as targets for antiviral drugs, e.g. dihydrofolate reductase gene can be inhibited with methotrexate. These can also be used to determine the immune status of a patient infected with HIV. HHV8 derived protein viral MIP II may be used as an anti-inflammatory agent for, e.g. treating rheumatoid arthritis. This sequence is stated as containing 81 open reading frames.

Sequence 137507 BP; 32579 A; 37795 C; 35758 G; 31375 T; 0 other;

Query Match 2.0%; Score 40.4; DB 19; Length 137507;
 Best Local Similarity 44.0%; Pred. No. 9.4;
 Matches 170; Conservative 0; Mismatches 216; Indels 0; Gaps 0;

```

Oy 514 CTGAAAAACAGATGAAGTCTCTGAGACGCGAGATGACCAACAGCTCGGAG 573
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 125106 CAGCAGATGAGCAGCAGCAGCAGATGAGCAGCAGCAGATGAGCAGCAGCAG 125047
Oy 574 GAGGCCACCGACATCAAGTGCAGTGAATGAAACATGAGAGCAATTTGAGCTCTCTCCAG 633
    ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| ||| |||
Db 125046 GATGACAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCAGCTTGAAG 124987
Oy 634 AGCCAGCGTCTGAGGTGAGAGAGATGATTCGAGACATGGGTGTGGACATGCGGTG 693

```

Db 124986 GAGCAGAGAGCAGAGGTTAGAGGATCAGAGAGAGTTAGAGGAGCAGAGCAGAGAGTTA 124927
Qy 694 GAGCAGCTGGCTGTCTACTGCTGCTCCCTCAGAAAGTATGAAATCTGAAAGAAAGCT 753
Db 124926 GAGAGAGCAGAGCAGAGGATTAGAGAGCAGAGCAGAGGATTAGAGAGCAGAGCAGAGAG 124867
Qy 754 CGAAGGCCACAGAGGAACTGGCTGACAGGTTGAAGAAGATTTGCTCCTTAGAGAGC 813
Db 124866 TTACAGAGAGCAGAGAGGATTAGAGAGCAGAGAGGATTAGAGAGCAGAGAGAGAG 124807
Qy 814 AAGTTGAAGACTCTCAACACTGAGCTGATCAGCCCAAGTTAGAACTGAGGTCAGCCAG 873
Db 124806 GAGTTAGAGAGCAGAGCAGAGGATTAGAGAGCAGAGGTTGGAAGCAAGAGCAGAGAG 124747
Qy 874 AAGGACTTACAAGTGTGAGCCAGGA 899
Db 124746 GTGGAAGAGCAAGAGCAGAGCAGGA 124721

RESULT 32
AAx83007/C
ID AAx83007 standard; DNA: 51259 BP.
XX
AC AAx83007;
XX
DT 31-AUG-1999 (first entry)
XX
DE Partial mouse WRN genomic sequence #3.
XX
KW Mouse; WRN; Werner's syndrome; detection; diagnosis; autosomal;
KM recessive disorder; phenotype; ss.
XX
OS Mus musculus.
XX
PN W09724435-A1.
XX
PD 10-JUL-1997.
XX
PF 30-DEC-1996; 96WO-US20785.
XX
PR 12-APR-1996; 96US-0632175.
PR 29-DEC-1995; 95US-0009409.
PR 29-DEC-1995; 95US-0580539.
PR 30-JAN-1996; 96US-0010835.
PR 30-JAN-1996; 96US-0594242.
XX
PA (DARW-) DARWIN MOLECULAR CORP.
PA (OSHI/) OSHIMA J.
PI Fu Y, Mulligan J, Oshima J, Schellenberg GP, Yu C;
DR WPI: 1997-363671/33.
XX
PT Isolated nucleic acid molecule encoding the WRN gene product -
PT useful for detection and treatment of Werner's syndrome, and related
PT diseases
XX
PS Claim 1; Fig 7; 153pp; English.
XX
CC This sequence represents a fragment of the genomic sequence containing
CC the coding region for the mouse WRN gene (AAx83004). The corresponding
CC human gene (AAx83001) encodes a protein related to Werner's syndrome.
CC The products can be used for the detection and treatment of Werner's
CC syndrome (WS), an autosomal recessive disorder with a complex phenotype,
CC as well as related diseases.
XX
SQ Sequence 51259 BP; 14533 A; 9635 C; 10266 G; 16825 T; 0 other;

Query Match 2.0%; Score 40.2; DB 18; Length 51259;
Best Local Similarity 48.1%; Pred. No. 6.3;
Matches 114; Conservative 0; Mismatches 123; Indels 0; Gaps 0;

Qy 513 CCTGAAAAACAGATGAAGTTCCTGAGCAGCGCAGAGATGACCAACAAGCTCGGA 572
Db 269 CATGAAG 210
Qy 573 GGAGGCCACCGACTCAAGTCAGAGTGAAGAAACATGAGAGCAAAATGACTCTACTCA 632
Db 209 GGAGGAGCA 150
Qy 633 GAGCCAGGCTTCTAGGTGAGAGAGATGATTCGACACTGGGTGTGGACAGTACCGGT 692
Db 149 GGAG 90
Qy 693 GAGCAGCTGCTGTCTGTACTGCTGCTGCTCCCTCAGAAAGATGAGAAATCTGAAGA 749
Db 89 GGAGGAG 33

RESULT 33
AAC59944
ID AAC59944 standard; cDNA: 941 BP.
XX
AC AAC59944;
XX
DT 30-JAN-2001 (first entry)
XX
DE Human secreted protein cDNA sequence #38.
XX
KW Cytostatic; immunosuppressive; neutropic; neuroprotective; antiviral;
KW antiallergic; hepatotropic; antidiabetic; antiinflammatory; antitumor;
KW vulnerrary; anticonvulsant; antibacterial; antifungal; antiparasitic;
KW cardiant; gene therapy; cancer; immune disorder; cardiovascular disorder;
KW neurological disease; infection; human; secreted protein; ss.
XX
KW Homo sapiens.
XX
OS
XX
PN W020005198-A1.
XX
PD 21-SEP-2000.
XX
PF 09-MAR-2000; 2000WO-US06012.
XX
PR 12-MAR-1999; 99US-0124093.
PR 23-NOV-1999; 99US-0166989.
XX
XX
PA (HUMA-) HUMAN GENOME SCI INC.
XX
PI Rosen CA, Ruben SM, Komatsoulis G;
DR WPI: 2000-587520/55.
DR P-PSDB: AAB28739.
XX
PT Human secreted proteins and the nucleic acids that encode them, useful
PT in gene therapy protocols and recombinant nucleic acid based procedures
XX
PS Claim 1; Page 337; 391pp; English.
XX
CC The invention relate to the isolation of genes AAC59907-C59956 encoding
CC 50 human secreted proteins AAB28702-B28751. The genes can be used to
CC generate fusion proteins by linking to the gene for the human
CC immunoglobulin G Fc portion for increasing the stability of
CC the fusion protein as compared to the human protein only. The genes and
CC proteins are useful for preventing, ameliorating or treating medical
CC conditions, e.g. by protein or gene therapy. The genes are isolated
CC from a range of human tissues disclosed in the specification. The
CC nucleic acids, proteins, antibodies and (ant)agonists are useful in
CC the diagnosis, treatment and prevention of: (a) cancer, e.g. breast
CC and ovarian cancer, and other cancers of the adrenal gland, bone, bone
CC marrow, breast, gastrointestinal tract, liver, lung, or urogenital;
CC (b) immune disorders e.g. Addison's disease, allergies, autoimmune
CC haemolytic anaemia, autoimmune thyroiditis, diabetes mellitus, Crohn's
CC disease, multiple sclerosis, rheumatoid arthritis and ulcerative

KX		pharmaceutical; gene; ss.
XX		
OS	Drosophila melanogaster.	
PV	WO200171042-A2.	
PN		
PD	27-SEP-2001.	
PF		
PA	(PERE) PE CORP NY.	
PI	Venter JC, Adams M, Li PWD, Myers EW;	
DR	WPT: 2001-656860/75. P-PSTB: ABB62220.	
PR	New isolated nucleic acid detection reagent for detecting 1000 or more genes from Drosophila and for elucidating cell signalling and cell-cell interactions -	
PS	Claim 1: SEQ ID NO 13451; 21pp + Sequence Listing; English.	
CC	The invention relates to an isolated nucleic acid detection reagent capable of detecting 1000 or more genes from Drosophila. The invention is useful in developmental biology and in elucidating cell signalling and cell-cell interactions in higher eukaryotes for the development of insecticides, therapeutics and pharmaceutical drugs. The invention discloses genomic DNA sequences (AB101840-AB16175) expressed DNA sequences (AB101840-AB16175) and the encoded proteins (ABB57737-ABB72072).	
CC	The sequence data for this patent did not form part of the printed specification, but was obtained in electronic format directly from WIPO at ftp://wipo.int/pub/published_pcr_sequences.	
SQ	Sequence 1931 BP; 543 A; 499 C; 552 G; 337 T; 0 other:	
Query Match	2.0%; Score 39.8; DB 23; Length 1931; Best Local Similarity 46.8%; Pred.No.1.4; Matches 125; Conservative 0; Mismatches 142; Indels 0; Gaps 0	
OY	476 AGACGCGCTTAACAAGCAGATGTGGTTCACCCTGAANAACAGTAGAATTGCC Db agaagaaccagaaacagaaaacagaaacagaagaacaagtatcgcacaagaagaacgaagg	535
Oy	536 TGGCACGACGCGCAGCATGTGACCAAAACAGCTCGGGAGAGAGCCCACGCATCAACTGCA Db gacacgacgcaagaagaaacgcaaacgcaacggcttaaatgtacgagcacgcttgctaagaacaa	595
OY	596 AGATAAAAACCATGAGAACAAATTGAGCTCTCTCCAGAGCCAGCCTTGAGAGTTGAGG DB agaaaaagcccgagcagcgcagataaagattcagaaaaaaaaagcagcataagtcataaaga	655
DB	1108 agaaaaagcccgagcagcgcagataaagattcagaaaaaaaaagcagcataagtcataaaga OY AGATGATTTCGACACATGGGTGTGGCACAGTCAAGCGGTGACAGCTGGCTGTCTACTGCG	715
DB	1168 agaaagaaccaagcagaagaatatccaatalcagcagagagatgaacaacagcttactcccgtlgaag OY 716 TGTCCCTCAGAAGAAAGTATGAGAAATC DB tgatcaaagaagccagtygctatlgaac	1227
RESULT	38	
ID	ABL06322 standard; cDNA; 3931 BP.	
XX	ABL06322;	
XT	26-MAR-2002 (first entry)	

xx	Drosophila melanogaster expressed polynucleotide SEQ ID NO 13448.
DE	
xx	Drosophila: developmental biology; cell signalling; insecticide;
KW	Pharmaceutical; gene; ss.
xx	
OS	Drosophila melanogaster.
xx	
PN	WO200171042-A2.
xx	
PD	27-SEP-2001.
xx	
PE	23-MAR-2001; 2001WO-US09231.
xx	
PR	23-MAR-2000; 2000US-191637P.
xx	11-JUL-2000; 2000US-0614150.
PA	(PEKE) PE CORP NY.
PI	Venter JC, Adams M, Li PWD, Myers EW;
xx	
DR	WPI: 2001-656860/75.
xx	P-PsDB; ABB62219.
PT	New isolated nucleic acid detection reagent for detecting 1000 or more
pt	genes from Drosophila and for elucidating cell signalling and cell-cell
xx	interactions -
PS	Claim 1: SEQ ID NO 13448; 21pp + Sequence Listing; English.
xx	
CC	The invention relates to an isolated nucleic acid detection reagent
CC	capable of detecting 1000 or more genes from Drosophila. The invention is
CC	useful in developmental biology and in elucidating cell signalling and
CC	cell-cell interactions in higher eukaryotes for the development of
CC	insecticides, therapeutics and pharmaceutical drugs. The invention
CC	discloses genomic DNA sequences (ABL16176-ABL30511), expressed DNA
CC	sequences (ABL01840-ABL16175) and the encoded proteins
CC	(ABB57737-ABB72072).
CC	The sequence data for this patent did not form part of the printed
CC	specification, but was obtained in electronic format directly from WIPO
CC	at ftp.wipo.int/pub/published_pct_sequences.
xx	
SQ	Sequence 3931 BP; 1137 A; 887 C; 966 G; 941 T; 0 other:
Query Match 2.0%; Score 39.8; DB 23; Length 3931;	
Best Local Similarity 46.8%; Pred. No. 2;	
Matches 125; Conservative 0; Mismatches 142; Indels 0; Gaps 0;	
OY	476 AGACGCCCTTAAACAAGCAGATGCTGTTCACCCCTGAATAAACAGATGAAGTTCC 535
Db	
1988	aagaagcaacagaaacaagaacacagaagaacgaagaatacagaacaagaagaacaagc 2047
OY	536 TGGACACCGGAGCATGTGACCACCAACAGCTCGGAGAGGCCCCACGCATCAAGTGCA 595
Db	
2048	gaccacgcacgagaagaacgcagaacacagggcccaaagacgaagcagctcgcaagcaaa 2107
OY	596 AGATGAAAACCATGTGAGCAAAATTGACTCTAATCTACAGACCGACGCTTCTGAGGTGAGC 655
Db	
2108	agaaaagcgcgagcagcagcagataaagtacgaanaagaagaagcagataaagtaacaaaga 2167
OY	656 AGATGATTTCGAGACATGGGTGTGGGACACACTCACGCCGTGGAGCAGCTGGCTGTGTAAGTGGC 715
Db	
2168	agaagaagcgcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagcagc 2227
OY	716 TGTCCTTCACAGAAGAGATGATGAGAATC 742
Db	
2228	tgatcaagaagccagtggtactatgtagac 2254
RESULT	39
ABA58819/c	
ID	ABA58819 standard; DNA: 475 BP.

XX ABA58819;
 AC
 XX 01-FEB-2002 (first entry)
 DT
 XX Human foetal liver single exon nucleic acid probe #7124.
 DE
 XX Human; foetal liver; gene expression; single exon nucleic acid probe; ss
 KW
 XX Homo sapiens.
 OS
 XX MO200157277-A2.
 PN
 XX 09-AUG-2001.
 PD
 XX 30-JAN-2001; 2001MO-US00669.
 PF
 XX 04-FEB-2000; 2000US-0180312.
 PR 26-MAY-2000; 2000US-0207456.
 PR 30-JUN-2000; 2000US-0608408.
 PR 03-AUG-2000; 2000US-0632366.
 PR 21-SEP-2000; 2000US-0234687.
 PR 27-SEP-2000; 2000US-0236359.
 PR 04-OCT-2000; 2000GB-0024263.
 XX
 PA (MOLE-) MOLECULAR DYNAMICS INC.
 XX
 PI Penn SG, Hanzel DK, Chen W, Rank DR;
 XX
 DR WPI; 2001-483447/52.
 XX
 PT Human genome-derived single exon nucleic acid probes useful for
 PT analyzing gene expression in human foetal liver -
 XX
 XX Claim 1; SEQ ID NO 7124; 639pp + sequence listing; English.
 XX
 CC The invention relates to a single exon nucleic acid probe for
 CC measuring human gene expression in a sample derived from human foetal
 CC liver. The single exon nucleic acid probes may be used for predicting
 CC measuring and displaying gene expression in samples derived from human
 CC foetal liver. The present sequence is a single exon nucleic acid
 CC probe of the invention.
 CC Note: The sequence data for this patent did not form part of the
 CC printed specification, but was obtained in electronic format directly
 CC from WIPO at ftp.wipo.int/pub/published_pat_sequences.
 XX
 XX Sequence 475 BP; 38 A; 198 C; 45 G; 194 T; 0 other;
 XX

Query Match	2.0%:	Score 39.6:	DB 22:	Length 475:
Best Local Similarity	46.4%:	Pred. No. 0.73:		
Matches 129:	Conservative	0:	Mismatches 149:	Indels 0:
				Gaps 0:
QY	483 CTTAAACAAGCCAGAGATCTGTTCCACCCCTGAAAAAACGATGAATTCCTGAGCA	542		
DB	401 CTTACGCCCTGGGAGACAGACAAGATCTGTCTCAAAAAACAGAAAGAGAGAGA	342		
QY	543 GCGGAGAGATGAGACCCAAACAGCTCGGAGAGAGGCCCAACCACTCAATGCAAGATGA	602		
DB	341 GAAGGAGGAAGGAGAGAGAGAGAGAAAGGAGAAAGAGAGACAGAGAGAAGAGA	282		
QY	603 AACCATGAGCAAAATTGAGCTCTACTCCAGAGCCAGCTTTGTGAGGTGGAGAGATGAT	662		
DB	281 GAAGGAAGGAAGGAAGAAAGAAAGAGAGAGAGAGAGAGAGAGAGAGAGAGA	222		
QY	663 TCGAGACAGTGGGTGTGGGACACTCAGCCGTTGGAGACAGCTGCTGTACTGCTGTCCCT	722		
DB	221 GAAGAGAAGGAAGAAAGAAAGAGAGAGAAAGAGAGAGAGAGAGAGAGAGAGA	162		
QY	723 CAAGAAAGAGATGAGAAATCTGAAGGAACCTCGGAAG	760		
DB	161 GAAGGAAG	124		

RESULT 40
 ABA27737/c
 ID ABA27737 standard; DNA; 475 BP.
 XX
 AC ABA27737;
 XX
 DT 23-JAN-2002 (first entry)
 XX
 DE Probe #6203 for gene expression analysis in human heart cell sample.
 XX
 KW Human; gene expression; heart; microarray; vascular system; probe;
 KW cardiovascular disease; hypertension; cardiac arrhythmia;
 KW congenital heart disease; ss.
 XX
 OS Homo sapiens.
 XX
 PN M0200157274-A2.
 XX
 PD 09-AUG-2001.
 XX
 PF 30-JAN-2001; 2001MO-US00666.
 XX
 PR 04-FEB-2000; 2000US-0180312.
 PR 26-MAY-2000; 2000US-0207456.
 PR 30-JUN-2000; 2000US-0608408.
 PR 03-AUG-2000; 2000US-0632366.
 PR 21-SEP-2000; 2000US-0234687.
 PR 27-SEP-2000; 2000US-0236559.
 PR 04-OCT-2000; 2000GB-0024263.
 XX
 PA (MOLE-) MOLECULAR DYNAMICS INC.
 XX
 PI Penn SG, Hanzel DK, Chen W, Rank DR;
 XX
 DR WPI; 2001-488899/53.
 XX
 PT Single exon nucleic acid probes for analyzing gene expression in human
 PT hearts -
 XX
 PS Claim 1; SEQ ID No 6203; 530pp; English.
 XX
 CC The present invention relates to single exon nucleic acid probes for
 CC measuring human gene expression in a sample derived from human heart. These
 CC present sequence is one such probe. The probes may be used for
 CC predicting, measuring and displaying gene expression in samples derived
 CC from the human heart via microarrays. By measuring gene expression, the
 CC probes are useful for predicting, diagnosing, grading, staging,
 CC monitoring and prognosing diseases of the human heart and vascular system
 CC e.g. cardiovascular disease, hypertension, cardiac arrhythmias and
 CC congenital heart disease.
 CC Note: The sequence data for this patent did not form part of the printed
 CC specification, but was obtained in electronic format directly from WIPO
 CC at ftp.wipo.int/pub/published_pat_sequences.
 XX
 SEQ Sequence 475 BP; 38 A; 198 C; 45 G; 194 T; 0 other;

Query Match	Similarity	Score	DB	Length
Best Local	46.4%	Pred. No. 0.73		
Matches	129	Conservative	0	Mismatches 149; Indels 0; Gaps 0;
Qy	483	CTTTAAACAAGCAGAGATCTGTGTTCCACCCGTAAGAAAACGATGAATTCTCGAGCA	542	
Db	401	CTTTCAGCCTGGAGACAGCAGATCCTGTCTCAAAAACGAAAGAACGAGAGAAGAA	342	
Qy	543	CGCGCAGATGTAGACCAAAACAGCTCGGAGGAGGCGCCACCACTCAATGCACAGATGAA	602	
Db	341	GAAAGGAGAAAGAGAGAGAGAGAGAAAGGAGAAAGGAGAAAGAGAGAGAGAGAGAA	282	
Qy	603	AACCATGAGACAATTGAGCTCTACTCCAGAGCCAGCGTCTGAGAGTGAGAGATGAT	662	
Db	281	GAAAGAA	222	

Oy 663 TCGAGACATGGGTGTGGGACAGTCAAGCGTGAGCAGCTGCTGTGTACTGCGTCCCT 722
| | | | | | | | | | | | | | | | | | | | | |
Db 221 GAAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGA 162
| | | | | | | | | | | | | | | | | | | | | |
Oy 723 CAAGAAGAGTATGAGAACTCTGAGGAAGCTCGAAGG 760
| | | | | | | | | | | | | | | | | | | | | |
Db 161 GAAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGAGAAGG 124
| | | | | | | | | | | | | | | | | | | | | |

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Job time: 9260 sec